Molecular functional analyses of larval adhesion in a highly fouling invasive model ascidian

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Received: 29 May 2022 / Accepted: 20 August 2022
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Abstract
Successful papillary adhesion is the first essential step for serious ascidian biofouling in marine ecosystems; however, the mechanisms of papillary adhesion have not been fully elucidated. Here, we employed transcriptome sequencing to assess the differentially expressed genes (DEGs) between papillae and remaining body in the larvae of the highly fouling invasive model ascidian Ciona robusta. Enrichment results for DEGs showed that papillae were adhesive structures with the combined functions of substrate recognition, environmental perception, and adhesive protein synthesis and secretion. Two candidate ascidian papilla adhesive proteins (APAPs), APAP-1 and APAP-2, were identified, expressed, and purified in vitro. Surface coating tests showed that APAP-1 was a cohesive protein while APAP-2 was an interfacial protein involved in the adhesion between papillae and material surface. Collectively, the obtained DEGs and adhesive proteins provide candidates to deeply understand molecular mechanisms of underwater adhesion and further develop anti-fouling strategies in marine ecosystems.

Keywords  Ascidians · Papillae · Biofouling · Adhesion · Transcriptome · Adhesive protein

Introduction

Biofouling, the intensive colonization and adhesion of organisms on submerged substrates, has caused a series of economic and environmental issues in marine ecosystems (Salta et al. 2010; Madin and Ching 2015; Bannister et al. 2019). So far, the number of marine fouling species has reached >4000, among which ascidians are one of the dominant taxa due to their enormous adhesion biomass on the surfaces of underwater facilities, causing significantly negative effects on both industries and local environments (Aldred and Clare 2014). For instance, fouling ascidians were the main contributor to the production loss of cultured sea cucumbers and shellfish such as oysters, clams, and scallops in the Pacific, Atlantic, and Indian Oceans, and the economic loss for the control of fouling species such as ascidians in aquaculture alone was approximately US $1.5—3 billion per year (Lambert 2007; Adams 2011; Fitridge et al. 2012). In addition, ascidian biofouling was also one of the additional costs for hull maintenance and fuel consumption in the shipping industry (Aldred and Clare 2014). Ecologically, fouling ascidians can reduce species richness and change biodiversity patterns of benthic communities in local marine ecosystems (Castilla et al. 2004; Zhan et al. 2015). For example, biofouling by the ascidian Ciona intestinalis in San Francisco Bay resulted in sharp decreases in biomass of other sessile species such as Bryozoa and Porifera (Blum et al. 2007). Although a series of encouraging progress have been made in anti-fouling in the past decades (Salta et al. 2010; de With 2018), unfortunately ascidian fouling has not been effectively controlled, which can be seen from an indisputable fact that the negative influence has been increasing in frequency and area affected (Aldred and Clare 2014; Zhan et al. 2015; Simkanin et al. 2016). Therefore, there is an urgent need to develop effective anti-fouling strategies to maintain the health and integrity of global marine ecosystems and sustainable development of marine industries.

The key step in formulating anti-fouling strategies is to successfully reveal the detailed mechanisms underlying rapid and strong adhesion to underwater surfaces...
(Prendergast 2009). Ascidian adhesion begins with swimming larvae and becomes stronger in adults. Solitary adult ascidians with high fouling abilities rely mainly on their special adhesive organ termed stolon to adhere firmly to various underwater surfaces (Hirose and Akahori 2004; Ueki et al. 2018). Studies have showed that ascidian stolon was insensitive to the changes of marine environments (Li et al. 2021a), thus increasing great uncertainty and difficulty to eradication. The adhesion of swimming larvae is a signal to initiate subsequent metamorphic development of ascidians (Matsunobu and Sasakura 2015). As a result, if larval adhesion fails, ascidians are not able to start metamorphosis to complete their life history, and finally die off. Therefore, when comparing with high difficulties in control and management of fouling ascidians at the adult stage, the interruption of adhesion at larval stages represents a direct, cost-effective, and efficient way to solving biofouling problems by ascidians.

The adhesion of ascidian larvae depends on their specific structures at the anterior end of the head, which are composed of three cup-shaped translucent papillae. Each papilla consists of three distinct types of cells: axial columnar cells (ACCs), lateral primary sensory neurons (PSNs), and central collocytes (CCs). ACCs and PSNs play important roles in substrate recognition, while CCs are involved in the synthesis of adhesive proteins, which are eventually transported to the hyaline caps prior to the papilla (Zeng et al. 2019a). Available evidence suggests that the adhesive proteins synthesized in the CCs of the papilla can be quickly released to substrate surfaces to achieve underwater adhesion when papillae successfully contact with the surfaces of proper underwater substrates (Cloney 1977; Pennati et al. 2007; Zeng et al. 2019b). However, in contrast to the detailed knowledge of structure of ascidian papillae, little is known about the protein composition of these papillary secretions and no associated adhesive protein has been successfully identified from papillae. The underlying molecular mechanisms of these protein adhesions, which are essential for formulating anti-fouling strategies against ascidian larvae, are also largely obscure. Indeed, studies have been trying to dissect adhesive mechanisms of marine fouling organisms, for example, a series of mussel adhesive proteins including collagens (pre-Cols, TMP and PTMP) and foot proteins (Mfp-1 ~ Mfp-6) have been successfully isolated and identified from marine mussel byssus. The foot protein Mfp-1 is distributed on the cuticle of byssus and plays a protective role on byssus structure. Mfp-2 and Mfp-4 are located in the core of plaque, which are connected to byssal thread. Mfp-3, Mfp-5, and Mfp-6 are located in the bottom of plaque and participate in the interface adhesion between byssus and underwater substrate surface (Hwang et al. 2010; Suhre et al. 2014; Waite 2017). All these results provide valuable reference for the functional analysis of adhesive proteins in ascidian larvae.

One of the main reasons for the lack of adhesive protein information is the small size of larval papillae (only 20 μm in length), largely impeding the isolation and identification of adhesive proteins from its tiny sub-structures. The advanced transcriptome sequencing technique developed recently can help solve this technical problem because it allows for the use of total RNA isolated from micro-samples for library construction and sequencing (Cao et al. 2019). Using this technique, the cDNA libraries of two micro-samples, the papillae and the remaining body of Ciona robusta larvae, were constructed in this study. C. robusta was used as the experimental model here mainly due to its strong fouling characteristics at both larvae and adult stages in marine ecosystems (Aldred and Clare 2014). In particular, this species is highly invasive, and the strong fouling capacity has caused severe economic and ecological problems globally (see review by Zhan et al. 2015). In contrast to its negative effects, the well sequenced and annotated small genome (123 Mb), as well as abundant genetic resources, makes this species a good model for answering multi-disciplinary questions (Zhan et al. 2015; Satou et al. 2019), thus facilitating detailed gene expression and protein function analyses in this study. After comparative transcriptomic analyses, we characterized differentially expressed genes (DEGs) between the two micro-samples, and subsequently genes encoding for candidate papilla adhesive proteins were identified from papillae and then expressed in vitro. The adhesive abilities of these purified proteins were further verified by surface coating analyses. Here, we aim to understand specific genes and proteins involved in papillary adhesion, and the results obtained in this study are expected to provide a reference to fully understand the mechanisms of underwater adhesion and develop anti-fouling strategies in marine ecosystems.

Materials and methods

Ascidian collection and artificial fertilization

Ciona robusta adults (6.0 cm in length) were collected from a marine shellfish farm in Dalian, Liaoning Province, China (38°49′13″ N, 121°24′20″ E) in September 2020. All collected ascidians were rapidly transported to the laboratory and temporarily acclimated in a 1000 L circulating aquarium using the same conditions (20 ± 1 °C, 30 ± 1 psu, and pH 8.10 ± 0.05) as the sampling site for three days. During acclimation, each hundred ascidians were fed daily with 10 g dried algae powder mixture of Chlorella sp. and Spirulina sp. until sperm duct and oviduct appeared. Six matured individuals were dissected with sterilized surgical blades to obtain sperm and eggs. The collected eggs were temporarily stored in a collection tube, while the collected sperm was added into a 1.5 mL centrifuge tube containing
1 mL pre-cooled sterilized seawater. Before fertilization, the eggs were rinsed with sterilized seawater for 1 min to remove their chorion surfaces. A total of 0.5 mL sperm was then added into the collection tube with 1000 eggs for artificial fertilization. After fertilization for 10 min, the eggs were rinsed slowly with sterilized seawater to remove excess sperm and incubated in a Petri dish. The incubation condition was controlled at 18°C ± 1, 30 ± 1 psu, and pH 8.10 ± 0.05 in darkness. The fertilized eggs were incubated for 24 h to obtain swimming larvae, which were finally collected for the following RNA extraction and transcriptome sequencing. The overall scheme of our experiment is shown in Fig. 1.

**Micro-dissection and RNA extraction**

The swimming larvae were transferred into 100 μL sterilized seawater dropped on a glass slide. A tungsten needle with tip diameter of 1 μm (HengMi, Shanghai, China) was used to dissect the papillae and body of *C. robusta* swimming larvae under an optical stereomicroscope (Olympus SZ51, Tokyo, Japan). The detailed dissecting position on swimming larvae is shown in Fig. 1. The dissected papillae and body were distributed into two 2 mL lyophilization tubes. Three biological replicates were set up for all the papillae and body samples, respectively. All samples were snap-frozen with liquid nitrogen and then preserved at −80°C.

The tunic on the body surface of ascidian larvae is impermeable for lysis solution when isolating the total RNA of larvae. Therefore, the collected samples were pre-fragmented using an ultrasonic cell disruptor (Scientz JY92-IIDN, Ningbo, China) with Φ II horn after being mixed with 350 μL cell lysis buffer from a RNeasy Micro Kit (Qiagen, 74,004, Dusseldorf, Germany). The whole ultrasound process was conducted on ice with parameters of 20 kHz, 60 W, and 2 min (1 s ON and 3 s OFF). The success of fragmentation and cleavage was approved by no large tissue fragments detected under a microscope. The total RNA was extracted from the disrupted papillae and body samples using a RNeasy Micro Kit following the manufacturer’s instructions. The possible DNA contamination was removed by DNase I provided by the kit.

**Transcriptome sequencing**

A total of six cDNA libraries were constructed based on the extracted RNA samples using the Smart-seq2 method (Picelli et al. 2014). A total of 5 ng RNA was used for reverse transcription and rRNA was depleted with PolyA-selection method. Fifteen cycles of PCR were performed during cDNA pre-amplification. The KAPA Hyper Prep
changes in eight DEGs were calculated following the 2−∆∆CT reference gene (Fujikawa et al. 2010), and the relative fold of β-actin was used as the primer information for these selected DEGs are listed in Table S1. The β-actin of C. robusta was used as the reference gene (Fujikawa et al. 2010), and the relative fold changes in eight DEGs were calculated following the 2−ΔΔCT method (Livak and Schmittgen 2001).

**Real-time quantitative PCR verification**

To verify the gene expression results obtained from transcriptome analysis, the same RNA samples used for transcriptome sequencing were also used to synthesize the first-strand cDNA using a reverse transcription kit (Takara, Tokyo, Japan) following the manufacturer’s instructions. In this experiment, a total of 40 ng of RNA for each sample was used for reverse transcription. A total of 5 up-regulated and 3 down-regulated DEGs were randomly selected from the transcriptome sequencing results to verify their expression changes by real-time quantitative PCR (RT-qPCR). The RT-qPCR was conducted using the LightCycler® 96 instrument (Roche, Basel, Switzerland) and a FastStart Essential DNA Green Master Kit (Roche) following the manufacturer’s instructions and the obtained 1 μL of first-strand cDNA was used as the PCR template. The detailed sequence and primer information for these selected DEGs are listed in the Table S1. The β-actin of C. robusta was used as the reference gene (Fujikawa et al. 2010), and the relative fold changes in eight DEGs were calculated following the 2−ΔΔCT method (Livak and Schmittgen 2001).

**Adhesion-related gene screening**

The genes encoding papilla adhesion-related proteins were screened from the DEGs with the fold changes of genes expression > 4 between papillae and body. The remaining genes were further screened according to the common characteristics of adhesive proteins in marine organisms summarized by previous literature (Li et al. 2021b). Briefly, the most abundant amino acids were glycine, leucine, tyrosine, serine, threonine, cysteine, or arginine and the percentages were greater than 10%. Meanwhile, the conserved domains such as thrombospondin-1 (TSP-1), von Willebrand type C (vWC), or epidermal growth factor (EGF) must be included. In addition, repetitive sequence and posttranslational modification (phosphorylation, glycosylation, and hydroxylation) were considered as optional screening criteria. The ProtParam and ProtScale online programs in Expasy platform (https://www.expasy.org/) were used to predict the molecular size, charge, and hydrophobicity of candidate adhesive proteins deduced from the screened genes. SMART (http://smart.embl-heidelberg.de/), SignalP (http://www.cbs.dtu.dk/services/SignalP), NetPhos-3.1 (https://services.healthtech.dtu.dk/service.php?NetPhos-3.1), SOPMA (https://npsa-prabi.ibcp.fr/cgi-bin/npsa_automat.pl?page=npsa_sopma.html), and PHYRE2 (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index) were employed to predict the conserved domains, signal peptides, phosphorylation sites, and spatial structure of candidate adhesive proteins, respectively.

**Protein recombination and purification**

The full-length open reading frames (ORFs) of the genes encoding for candidate papilla adhesive proteins were synthesized in vitro. The synthesized sequences were inserted into an expression plasmid pCMV3-C-His Negative Control Vector with a C-terminal decahistidine (10×His) tag (SinoBiological, Beijing, China). The recombinant vectors were then transformed into the mammalian cell HEK293 expression system to produce recombinant proteins. The recombinant HEK293 cells were sub-cultured for 7 days at 37°C and 5% of carbon dioxide, and the medium was exchanged every 2 days. A total of 10 mL of culture solution with a cell density of 4×10⁶ cells/mL was centrifuged for cell precipitation at 1000 rpm. The cells were lysed by 20 mL of ice-cold-modified RIPA lysis buffer with cocktail of protease inhibitors (Sigma, Burlington, USA). The lysis system was then filtered by the membrane with the diameter of 0.45 μm. The Ni affinity chromatography columns (GE 17–0409-01, Uppsala, Sweden) were used for adsorbing proteins from filtrate and the flow rate was 1 mL/min, after which the target proteins were gathered by gradient elution with imidazole. The concentrations of imidazole were 10, 50, 100, 200, 300, and 400 mM, and the flow rate was 2 mL/min. The protein concentration of eluent was detected by
the sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) method, and the recombinant protein was finally dissolved in the PBS after dialysis with 1×PBS buffer. The concentration and purity of adhesive proteins were determined by UV spectroscopy with optical density value at 280 nm and SDS-PAGE method, respectively.

**Adhesive ability test**

Surface coating experiment was carried out on different materials to test the adhesive abilities of recombinant proteins according to a previous method with minor modifications (Li et al. 2019). Polyvinyl chloride (PVC), acrylonitrile–butadiene–styrene plastic (ABS), mica, stainless steel, aluminum, hydrophilic glass, and hydrophobic glass were selected as the substrate materials. The concentration of all recombinant proteins was adjusted to 0.22 mg/mL before tests and the test volume was 10 μL. The Cell-Tak (Corning Life Sciences, New York, MA, USA), a commercial adhesive mixture of mussel foot proteins, was used as the positive control, while the bovine serum albumin (BSA, Yuanye, Shanghai, China) was used as the negative control. The PVC material was further used to estimate the effects of metal ions (calcium, magnesium, sodium, and potassium ions) and charged amino acid (glutamate and arginine) on the adhesive abilities of recombinant proteins. For all tests, the spots of proteins were stained using Bradford assay (Solarbio, Beijing, China) after dropping on the material surfaces for 12 h. The stained spots were photographed with a digital camera (E-P7, Olympus, Tokyo, Japan) and obtained images were submitted to the ImageJ software (Hartig 2013) for gray analysis. The peak plot and relative gray value were used to represent the adhesive abilities of recombinant proteins.

**Statistical analysis**

The difference between the relative gray values of spots in adhesive ability analysis was evaluated using one-way analysis of variance (ANOVA) with the significance levels at \(*p < 0.05\) and \(**p < 0.01\). The gene expression changes from DEGs analysis were represented by volcano plot. The significantly enriched functional clusters in GO and KEGG analyses were selected for plotting using ggplot2 in the R package (Wickham et al. 2016). The rest of the plots were drawn using Adobe Illustrator CS5.

**Results**

**Transcriptome sequencing**

The quality of the extracted RNA from the sample treated with ultrasonication met the standard of cDNA library construction (Fig. S1). On average, 28.38 million paired-end reads were produced by sequencing the cDNA libraries constructed from the papillae and body samples. After adapter cleaning, quality trimming, and length filtering, a total of 28.00 million clean reads (98.67% of the total reads) were obtained. For the papillae cDNA library, an average of 62.75% of the quality-filtered reads were mapped to the reference genome of *C. robusta* and 2.85% of the reads were mapped to multiple regions of this genome. For the body cDNA library, the correspondence mapping ratios were 75.47% and 3.61%, respectively (Table S2). These multiple mapping reads were excluded in the subsequent analysis by HiSAT2. A total of 18,191 genes were annotated into the genome of the ascidian *C. robusta* (Satou et al. 2019), and an average of 17,633 (±173) genes were found to be shared in six samples.

**Identification of DEGs**

After filtering the genes with low read counts, a total of 16,001 genes were obtained for subsequent DEGs analysis. A total of 5,051 genes were found to be differentially expressed in both papillae and body samples, with 1,875 genes were highly expressed in papillae and 3,176 genes were highly expressed in body (Fig. S2; Table S3). The PCA result based on these DEGs illustrated that papillae and body samples were clustered separately (Fig. S3). Analysis showed that the fold changes for the DEGs in RT-qPCR and transcriptome sequencing had good correlations, confirming the accuracy and validity of our transcriptome sequencing results (Fig. S4).

**Functional enrichment of DEGs**

A total of 1875 genes highly expressed in papillae compared with body were arranged for GO and KEGG enrichments. A total of 174 terms were obtained from GO enrichment and 20 terms were significantly enriched (Fig. 2a; Table S4). In the biological process classification, the DEGs were annotated to the terms ‘excretion’, ‘calcium ion transport’, ‘detection of external stimulus’, ‘face morphogenesis’, ‘protein dephosphorylation’, ‘aminoglycan catabolic process’, and ‘multivesicular body organization’. In the cellular component classification, the DEGs were enriched into the terms ‘basement membrane’, ‘photoreceptor outer segment’, ‘external encapsulating structure’, ‘pseudopodium’, ‘non-motile cilium’, and ‘neuron projection membrane’. In addition, the terms ‘serine hydrolase activity’, ‘transmission system’, ‘iron ion’, and ‘proteoglycan and glycosaminoglycan binding’ were enriched into the classification of molecular function. Taken together, the enriched GO terms could be divided into four categories including neural recognition, environmental perception, metabolism, and ion transport.
The KEGG enrichment results could also be divided into four categories: neural recognition, environmental perception, metabolism, and ion transport (Fig. 2b, Table S5). The pathways ‘serotonergic synapse’ and ‘neuroactive ligand-receptor interaction’ were enriched in neural recognition category. The pathways ‘sensory system’ and ‘interaction of environmental information processing’ were associated with environmental perception. The metabolism category included ‘xenobiotics biodegradation’, ‘linoleic acid’, ‘purine’, and ‘nucleotide and arachidonic acid’ pathways. The ‘calcium signaling pathway’ was classified into the ion transport category. The KEGG pathways with higher values of rich factor were ‘non-homologous end-joining’, ‘linoleic acid metabolism’, and ‘ovarian steroidogenesis’ (Table S5).

**Genes involved in papillary adhesion**

A total of six genes, encoding candidate papilla adhesive proteins with molecular size ranging from 18.54 kDa to 116.29 kDa were identified (Table 1). The proteins encoded by these genes were typically characterized by a high proportion of serine, cysteine, glycine, and threonine, while most of these proteins had a proportion of hydrophobic amino acids such as leucine. Functionally conserved domains such as TSP-1, vWC, and EGF associated with the underwater adhesion of marine fouling organisms were distributed in the sequences of candidate papilla adhesive proteins. Some proteins were predicted to be of repetitive sequences, including TSP-1 and LDLa (cysteine-rich) repetitive sequences in ‘SCO-spondin’ (Gene id: KY.Chr1.200.
v3.ND3-3), Fibronectin type 1 (FN1) repeat sequence in ‘Chordin-like protein 1’ (Gene id: KY.Chr1.10268. v1.ND1-1), and several internal repeats in ‘Protein delta homolog 2’ (Gene id: KY.Chr11.146.v1.SL1-1). In addition, some specific domains were also found in these candidate papilla adhesive proteins. The ‘Sushi, von Willebrand factor type A, EGF and pentraxin domain-containing protein 1’ (Gene id: KY.Chr10.1028.v1.nonSL5-1) possibly possessed an antimicrobial peptide-like domain, a protease activity inhibitory structural domain Somatomedin-B (SO) and a Bowman–Birk domain (BowB). Similarly, there was a leucine-rich SCAN domain in ‘Fibronectin’ (Gene id: KY.Chr2.475.v2.SL2-1), a cysteine-rich domain Defensin (DEFSN) and an ADAM Cysteine-Rich domain (ACR) in ‘Chordin-like protein 1’.

In addition to the genes directly associated with underwater adhesion, two genes, ‘opsin 1, long wave sensitive’ (OPN1LW, KY.Chr1.1186.v2.ND1-2) and ‘G protein-coupled receptor 84’ (GPR84, KY.Chr7.1150.v1.ND1-1), we screened on those that might be involved in neural perception in papillae (Table 1). Both proteins encoded by these genes had high percentage of leucine and carried 7 transmembrane receptor (rhodopsin family, 7tm_1) domains.

### Table 1

| Category                     | Gene ID                  | Gene description                                      | Molecular size (kDa) | Amino acid composition (top three) | Domain                                      | \(\log_2\) (fold change) | \(Q\) value |
|------------------------------|--------------------------|------------------------------------------------------|----------------------|-----------------------------------|---------------------------------------------|--------------------------|-------------|
| Papilla adhesive proteins    | KY.Chr1.200.v3.ND3-3     | SCO-spondin                                           | 116.29               | Ser12.8%, Cys11.5%, Thr7.1%       | TSP-1, vWC_def                              | 4.54                     | 7.01E-05    |
|                              | KY.Chr11.377.v2.ND2-1    | Spondin 2                                             | 18.54                | Gly12.9%, Lys11.1%, Ser9.9%       | TSP-1                                       | 7.58                     | 9.04E-05    |
|                              | KY.Chr2.475.v2.SL2-1     | Fibronectin                                           | 49.41                | Arg11%, Val8.2%, Thr7.5%          | FN1, FOLN, vWC                              | 3.16                     | 1.52E-04    |
|                              | KY.Chr11.146.v1.SL1-1    | Delta like non-canonical notch ligand 2               | 53.14                | Thr11.3%, Gly8.8%, Cys8.4%        | EGF, EGF_CA, EGF_like                       | 2.66                     | 1.61E-04    |
|                              | KY.Chr10.1268.v1.ND1-1   | Chordin-like                                          | 49.8                 | Ser12%, Cys7.9%, Arg7.4%          | vWC                                         | 2.07                     | 2.94E-04    |
|                              | KY.Chr10.1028.v1.nonSL5-1| Sushi, von Willebrand factor type A, EGF and pentraxin domain-containing 1 | 34.88 | Cys11.8%, Glu10.2%, Gly9.6% | EGF, EGF_CA, EGF_like, GLA | 3.52 | 2.24E-02 |
| Neural perception-related proteins | *KY.Chr1.1186.v2.ND1-2 * | Opsin 1, long wave sensitive                          | 44.36 | Leu10.4%, Val10.2%, Ile7.6% | Pfam:7tm_1 | 5.40 | 2.32E-05 |
|                              | *KY.Chr7.1150.v1.ND1-1   | G protein-coupled receptor 84                         | 31.06                | Ser10.3%, Lys9.6%, Leu8.5%        | Pfam:7tm_1                                  | 2.69                     | 8.11E-04    |

**Adhesive protein expression and purification**

A total of two proteins (Gene id: KY.Chr10.1028. v1.nonSL5-1 and KY.Chr10.1268.v1.ND1-1) were successfully expressed and purified in vitro. We named ascidian papilla adhesive protein 1 (APAP-1) and ascidian papilla adhesive protein 2 (APAP-2). After removing signal peptides and decahistidine tag labels, the purified product of recombinant APAP-1 had an SDS-PAGE band size of 46.47 kDa (Fig. 3a). The most abundant amino acid of this protein was cysteine, followed by aspartic acid, glycine, and asparagine in turn. Sequence analysis showed that APAP-1 had a large number of negatively charged amino acids (Fig. 3c). Furthermore, a total of six EGF-related domains, including one EGF, one EGF-like, and four EGF_CA domains, were found in APAP-1 (Fig. 3e). All the EGF-related domains were arranged in series with equal spacing, and there was only one amino acid spacing between two adjacent EGF-related domains. The purified product of recombinant APAP-2 had a band size of 65.01 kDa (Fig. 3b). The most abundant amino acid in this protein was serine, followed by cysteine, arginine, and lysine. Many amino acids in APAP-2 are phosphorylable, especially for serine (Fig. 3d). APAP-2 contained three vWC domains in tandem (Fig. 3f), which were distributed at the positions of 13–70 aa, 123–184 aa, and 210–270 aa of this protein sequences. It is predicted that both of APAP-1 and APAP-2 have the folding structures of extended strand, beta turn, and random coil (Fig. S5).

**Adhesive ability**

Surface coating experiment showed that all the positive controls (Cell-Tak) were stained obviously and the negative controls (BSA) were not stained (Fig. 4a). The APAP-1 on all the material surfaces were not stained, while the APAP-2 on the corresponding surfaces were stained obviously. The
adhesive abilities of APAP-2 on PVC, ABS, mica, stainless steel, aluminum, hydrophilic glass, and hydrophobic glass surfaces were strong, especially on aluminum surface where the color of APAP-2 spot was uniquely deeper than the corresponding positive control. There were no observed effects of metal ions on the adhesive ability of APAP-1 to PVC surface except iron. The additions of amino acids also had no interference with the adhesion of APAP-1 to PVC surface. (Fig. 4b). The iron also influenced the adhesive ability of APAP-2 on PVC surface (Fig. 4c). The changes in peak plots in all the gray analyses also showed similar trends with staining results (Fig. S6).

Discussion

In this study, we used transcriptome sequencing technique to assess the gene expression difference between the papillae and body tissues of *C. robusta* larvae. Interestingly, we obtained a large number of genes with higher expression levels in papillae when compared with body. Those identified genes were further functionally enriched into a series of terms/pathways, such as neural recognition, environmental perception, metabolism, and ion transport. Moreover, two adhesive proteins APAP-1 and APAP-2 were screened, expressed, and purified in vitro, and further confirmed to...
be associated with papillary adhesion. All these results provide essential information underlying molecular mechanisms of ascidian papillary adhesion, especially in the papillary substrate recognition, photoperception, and synthesis and secretion of adhesive proteins.

**Substrate recognition**

The larval adhesion of marine organisms to underwater substrates with different properties begins with their detective behaviors to substrate surfaces based on their nervous system (Prendergast 2009). In this study, functional enrichments for the up-regulated DEGs yielded many terms/pathways related to neural recognition, including ‘transmitter-gated channel activity’, ‘excitatory extracellular ligand-gated ion channel activity’, ‘neurotransmitter receptor activity involved in the regulation of postsynaptic membrane potential’, and ‘neuron projection membrane’, indicating the existing of strong neurophysiological activity in the papillae of *C. robusta*. In former studies, nervous system-like structures in papillae have been observed in the fouling ascidians including *C. intestinalis* (Imai and Meinertzhagen 2007a, b; Horie et al. 2009; Zeng et al. 2019b), *Distaplia occidentalis* (Cloney 1977), *Diplosoma macdonaldi* (Torrence and Cloney 1983), and...
Botryllus schlosseri (Caicci et al. 2010). These structures can be detected at the early tail-bud stage during embryo development using fluorescent labeling technique, and further were thought to be involved in ascidian larval settlement and metamorphosis (Imai and Meinertzhagen 2007b; Caicci et al. 2010). In consistent with findings in those studies, our results here showed the neural recognition behaviors in papillae at the molecular level, suggesting their importance for C. robusta larval adhesion.

Microvilli and cilia are common structures at the protrusions of neurofunctional cells in ascidian larvae (Hotta et al. 2007). Consistently, our results at the molecular level showed that DEGs were enriched in the terms of ‘non-motile cilium’ and ‘ciliary base’ (Fig. 2a, Table S4). These structures can be found in two kinds of cells, axial columnar cells (ACCs) and rostral trunk epidermal neurons (RTENs), with different morphologies and localizations in the papilla, both of which were thought to be of neural functions (Imai and Meinertzhagen 2007b; Caicci et al. 2010; Zeng et al. 2019b). During larval adhesion, ACCs in the papilla were functioned as mechanoreceptors contacting with substrate surfaces, while RTENs were considered as chemoreceptors that can primarily mediate metamorphic events after larval adhesion (Pennati et al. 2007). Such mechanical and chemical behaviors were observed in the larvae of ascidians Phallusia mammillata and D. macdonaldi (Torrence and Cloney 1983; Groppelli et al. 2003; Chase et al. 2016), as well as larvae of other marine adhesive organisms such as barnacles (Chaw and Birch 2009; Maruzzo et al. 2011). For example, the cyprids of barnacle had antennular setae, which were considered to be bimodal receptors with both chemo- and mechnano-receptive modalities, playing important roles in sensing hydrodynamic forces and dissolving substances and substrates (Maruzzo et al. 2011). Based on our transcriptome results, the swimming larvae of C. robusta may also have substrate recognition ability using these two types of cells.

Previous investigations demonstrated that light stimulus was an important environmental factor influencing the swimming behavior of larvae of marine organisms (Tsuda et al. 2003; Hirai et al. 2017). In fact, ascidian larvae had a specific pattern of swimming behaviors. In the first 3 h after hatching, they swim upward, and then changed to swim or sink downwards until settlement (Svane and Young 1989). However, it is difficult to establish the relationship between light stimulation and larval adhesion of marine organisms based on such evidence. In our results, the up-regulated DEGs were enriched in the terms/pathways of ‘detection of external stimulus’, ‘detection of light stimulus’, ‘photoreceptor outer segment’, and ‘sensory system’ (Fig. 2), suggesting that the neural structures in papillae of C. robusta should be involved in not only substrate recognition but also environmental perception, especially for photoperception during papillary adhesion. This finding is consistent with the result of an investigation on the larvae of the ascidian C. intestinalis, where the genes associated with chordate eye formation were highly expressed in papillae. Meanwhile, once the expressions of these genes were inhibited, the development of papillae was disturbed and the photosensitive swimming behavior was lost, influencing papillary adhesive ability (D’Aniello et al. 2006). In addition to ascidians, photoperception behaviors during larval adhesion have also been found in other aquatic organisms such as the fouling barnacle Balanus amphitrite (Kon-ya and Miki 1994) and mussel Dreissena polymorpha (Marsden and Landsky 2011). The results obtained here, as well as supportive evidence in related species, suggest that the success and firm underwater adhesion of C. robusta papillae should be closely related to their recognition to substrate surfaces stimulated by light.

Adhesive protein synthesis and secretion

Once ascidian papillae successfully recognize suitable submerged substrates, they begin to secrete adhesives with proteins as the main functional components. These adhesives were synthesized by the specific cells and then transported to hyaline cap, the site at the anterior end of each papilla (Cloney 1977). Our study obtained several up-regulated GO terms related to secretion functions, including ‘excretion’ and ‘multivesicular body organization’, which were in line with the fact that the massive adhesive proteins were synthesized during papillary adhesion in other ascidians (Torrence and Cloney 1983; Caicci et al. 2010). Collectively, papillae in C. robusta are an important structure responsible for the synthesis, storage, and secretion of adhesive proteins.

In the fouling ascidian C. intestinalis, the cells with secretory functions were localized at the trailing end of the papilla with microvilli, small endocytic vesicles, and large numbers of adhesive granules (Zeng et al. 2019b). Such secretory cells were also observed in the papilla of another fouling ascidian D. occidentalis (Cloney 1977). Transmission electron microscope analysis illustrated the presence of two types of collocyte granules with different electron dense in papillary secretory cells. These granules were rich in adhesive proteins and wrapped in circular or oval shape by some membrane structures, playing key roles in papillary adhesion in C. intestinalis by granule cross-linking (Zeng et al. 2019b). Importantly, genes associated with Ca^{2+} transport were found to be expressed in collocytes, suggesting that Ca^{2+} should be involved in the functioning of granules (Pang and Südhof 2010; Zeng et al. 2019b). Similarly, the relevant terms/pathways responsible for Ca^{2+} transport were enriched in our study (Fig. 2), suggesting that Ca^{2+} may act as an essential ion to participate in the synthesis, storage, and secretion of adhesive proteins in C. robusta papillae.
The effective composition of adhesives in marine fouling organisms has always been the focus in the relative research fields. In our study, the up-regulated proteoglycan-related terms such as ‘proteoglycan binding’, ‘glycosaminoglycan binding’, and ‘aminoglycan catabolic processes’ were significantly enriched in papillae (Fig. 2a). In adhesive organs, glycoproteins were one of the key biomacromolecules involved in interfacial adhesion between adhesive structures/tissues/organs and substrates (Sarosiek et al. 1988; Moussa et al. 2014; Opell et al. 2019). In particular, protein glycosylation has been detected in the adhesive structures of several marine fouling organisms, such as mussel byssus (Suhre et al. 2014) and barnacle cement (Kamino et al. 2012). Element and histochemistry analyses for the papillae of *C. intestinalis* showed that the adhesive granules in collocytes contained glycoproteins (Zeng et al. 2019a). In addition, carbohydrate-specific lectins could be detected in collocytes, hyaline caps, and adhesive plaques of the papillae in *C. intestinalis*, suggesting that the adhesives in larval papillae of this species contained some kinds of carbohydrate components, most likely in the form of proteoglycans (Zeng et al. 2019b). Aminoglycans, including hyaluronic acid and heparin, are important components of proteoglycans. In the study by Zeng et al. (2019a), the addition of heparin to seawater significantly reduced the adhesive rate of *C. intestinalis* larvae to the surface of plastic petri dish. The enrichments of the up-regulated GO terms ‘aminoglycan’, ‘glycosaminoglycan’, and ‘heparin’ in our study suggest that aminoglycans might be involved in the papillary adhesion of *C. robusta* (Table S4).

**Underwater adhesion of APAPs**

Two candidate proteins APAP-1 and APAP-2 involved in papillary adhesion in *C. robusta* were purified in this study. High abundance of cysteine, tandem EGF-related domain, and extremely negative electrical amino acids were the typical feature of APAP-1 sequence. The repeated cysteine residues and charged amino acids have been proved to be the important sequence structures that can regulate protein adhesion in marine fouling organisms. For example, CP-20k, an adhesive protein isolated from the barnacle cement, contained an arrangement of six repeated cysteine residues and negatively charged amino acids, which can form intramolecular disulfide bonds to act as a specific coupling agent between barnacles and calcified materials (Kamino 2001). The EGF domains are also involved in mediating protein–protein and protein–metal interactions in the adhesive organs/tissues in marine fouling organisms (Li et al. 2021b). Some adhesive proteins such as mussel foot protein Mfp-2 and sea star foot protein Sfp-1 were rich in EGF domains (Hwang et al. 2010; Hennebert et al. 2014). Mfp-2 was located in the adhesive plaque rather than the cuticle of mussel byssus, providing cohesion for the plaque to stabilize byssus structure. Atomic force microscope analysis showed that although Mfp-2 cannot directly adhere to the mica surface, there were strong interactions among these protein molecules by formatting bis- or tris-DOPA-ion complexes between the EGF domains of Mfp-2 and metal ions. The additions of Ca$^{2+}$ and Fe$^{3+}$ into the solution of this protein significantly enhanced these interactions, suggesting that the cohesiveness of adhesive plaque should rely on the ion complexation between EGF domains in Mfp-2 (Hwang et al. 2010). The EGF-related domains, including EGF, EGF-like, and four EGF_CA domains, account for 70% of the total protein theoretical size of APAP-1 in our study. In the surface coating experiment, the addition of Fe$^{3+}$ could significantly increase the adhesive ability of APAP-1 to PVC surface, even to the similar adhesive strength as the positive control (Fig. 4b). In addition, the GO term ‘iron ion binding’ was also significantly enriched in papillae by comparing the up-regulated DEGs between different samples. These results suggest that APAP-1 may be a cohesive protein involved in papillary adhesion in *C. robusta* and it requires the participation of some specific metal ions to promote the interactions between molecules of this protein (Fig. 5).

There were three vWF-like domains in the sequence of APAP-2 in *C. robusta*. vWF domain is a conserved structure distributed in many adhesive proteins in marine fouling organisms, such as PTMP-1 in mussels, TSP-1 containing protein in oysters, and Sfp-1 in sea stars (Hennebert et al. 2014; Suhre et al. 2014; Liu et al. 2016). Another attractive feature of APAP-2 is the high serine content in its amino acid composition. Similarly to APAP-2 in *C. robusta*, the cement proteins CP-19k and CP-68k in fouling barnacles and the foot proteins Mfp-5 and Mfp-6 in fouling mussels were also rich in serine (Kamino et al. 2000; Waite 2001; Zhao and Waite 2006; Urushida et al. 2007). In addition, the phosphorylation prediction for APAP-2 showed that most serine sites in this protein could be phosphorylated. The phosphorylated serine was abundance in both mussel foot proteins Mfp-5 and Mfp-6. The phosphorylated sites in Mfp-5 were associated with the conversion of serine to O-phosphoserine that can connect with the acidic mineral-binding motifs, contributing to the interfacial adhesion of mussel byssus to the calcareous substrates (Waite 2001; Zhao and Waite 2006). Phosphorylated proteins were also identified from the uncured cement of the fouling barnacle *Amphibalanus amphitrite*, which were further located in the organic matrix of base plate and capillary ducts, demonstrating that these phosphorylated proteins can form strong ionic bonds with underwater minerals and induce the mineralization of calcium carbonate (Dickinson et al. 2016). Cement was usually used to build the mineralized tubes of the sandcastle worm *Phragmatopoma californica*. Pc-3 was one of the three adhesive proteins with higher polarity in
worm cement. It was rich in serine residues (60–90 mol%) and most of which were phosphorylated (Zhao et al. 2005). It has been suggested that there were complex interactions among the phosphorylated serine of adhesive proteins, the free Ca$^{2+}$ in surrounding waters, and the submersed calcium substrates during the underwater adhesion of marine fouling organisms. A thrombospondin-containing byssal protein TSP-1 in the pearl oyster *Pinctada fucata* could aggregate and self-assemble through the interactions between Ca$^{2+}$ and phosphorylated serine or vWF domains in this protein, promoting the formation of byssus and binding ability of byssus to calcium substrates. In addition, the phosphorylation site of adhesive proteins was also supposed to be of strong electrostatic attraction with Ca$^{2+}$, forming the rigid structures of silks (Liu et al. 2016; George and Veis 2008). Overall, the sequence features of APAP-2 suggest that it may function as an adhesive protein taking part in the interfacial adhesion between the papillae of *C. robusta* and underwater substrates by regulating the interaction of Ca$^{2+}$, phosphorylated amino acids, and conserved domains. This conclusion was further supported by the evidence that the up-regulated terms/pathways related to Ca$^{2+}$ transport and serine phosphorylation were significantly enriched in papillae (Fig. 2). Importantly, the recombinant protein APAP-2 exhibits strong adhesive abilities to the surfaces of various materials in our surface coating experiments (Fig. 4a).

**Conclusion**

Using micro-dissection and transcriptome sequencing technologies, we obtained a gene pool that was highly expressed in papillae, the fouling organs of ascidian larvae. The terms/pathways related to neural recognition, environmental perception, metabolism, and ion transport were significantly enriched in functional analyses, confirming the important roles of these processes during the papillary adhesion of ascidian larvae. In addition, two candidate adhesive proteins APAP-1 and APAP-2 were identified, expressed, and purified in vitro, and functionally, they were proved to be a cohesive protein and interfacial protein, respectively, in the papillary adhesion. Multiple lines of evidence in our study clearly illustrate that papillae are adhesive structures in ascidian larvae with multiple functions of substrate recognition, environmental perception, and proteins’ synthesis and secretion.
Acknowledgements Great thanks to Profs. Zunchun Zhou and Bei Jiang for their assistance in ascidian collection. Great thanks to the Associate Editor, Dr. Mathias Wegner, and anonymous reviewers for their detailed, insightful, and constructive comments on early versions of this paper.

Author contributions Conceptualization: ABZ and SGL; methodology: JWC and XL; formal analysis: JWC and RYF; investigation: JWC and RYF; project administration: ABZ and SGL; funding acquisition: ABZ and SGL.

Funding This work was supported by the National Natural Science Foundation of China (Grant Nos. 42076098 and 32061143012), Youth Innovation Promotion Association, Chinese Academy of Sciences (Grant No. 20180854).

Data availability All the raw sequencing data of transcriptome sequencing were deposited in the National Centre for Biotechnology Information (NCBI) under the accession number (PRJNA795827).

Declarations Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval Not applicable.

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