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4	
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26 Abstract

Ability to tolerate low salinity is a key factor affecting the distribution of the Chinese 27 28 shrimp (Fenneropenaeus chinensis). Although previous studies have investigated the mechanisms underlying adaptations to low salinity in some crustaceans, little is 29 known about low-salinity adaptations in F. chinensis, particularly at the molecular 30 level. Here, to identify genes potentially associated with the molecular response of F. 31 chinensis to low-salinity exposure, we compared the transcriptomes of F. chinensis in 32 low-salinity (5 ppt) and normal-salinity (20 ppt) environments. In total, 45,297,936 33 34 and 44,685,728 clean reads were acquired from the low-salinity and control groups, respectively. De novo assembly of the clean reads yielded 159,130 unigenes, with an 35 average length of 662.82 bp. Of these unigenes, only a small fraction (10.5% on 36 37 average) were successfully annotated against six databases. We identified 3,658 differentially expressed genes (DEGs) between the low-salinity group and the control 38 group: 1,755 DEGs were downregulated in the low-salinity group as compared to the 39 40 control, and 1,903 were upregulated. Of these DEGs, 282 were significantly overrepresented in 38 KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways. 41 Notably, several DEGs were associated with pathways important for osmoregulation, 42 including the mineral absorption pathway (ATP1A, Sodium/potassium-transporting) 43 ATPase subunit alpha; CLCN2, Chloride channel 2; HMOX2, Heme oxygenase 2; 44 SLC40A1/FPN1, Solute carrier family 40 iron-regulated transporter, member 1), the 45 vasopressin-regulated water reabsorption pathway (AQP4, Aquaporin-4; VAMP2, 46 Vesicle-associated membrane protein 2; RAB5, Ras-related protein Rab-5) and the 47

ribosome pathway. Our results help to clarify the molecular basis of low-salinityadaptations in *F. chinensis*.

Key words: *ATP1A*; *AQP4*; Mineral absorption; Vasopressin-regulated water
reabsorption

52 Introduction

To improve the low-salinity aquaculture of marine crustaceans, it is necessary to understand their adaptation ability to low salinity and mechanisms used by these organisms to tolerate low-salinity environments. As osmoregulators, some euryhaline marine and brackish crustaceans have a strong ability to adapt to environments with varying salinities (from almost 0 ppt up to 40 ppt) [1-4]. This ability to tolerate low salinity is a key factor affecting the distribution of such crustaceans in low-salinity environments [5-6].

Previous studies have explored the mechanisms underlying low-salinity tolerance 60 in mariculture crustaceans at the organismal, cellular, and molecular levels [1, 7-10]. 61 62 In general, the most common adaptive strategies for hyperosmoregulation aim to maintain hemolymph osmolarity above that of the ambient medium, both via salt 63 absorption and via permeability reduction (i.e., reducing or limiting water inflow) [3]. 64 However, most euryhaline crustaceans produce isosmotic urine, and thus considerable 65 salt is lost in low-salinity environments [11-12]. Studies of low-salinity tolerance in 66 crustaceans have shown that the gills also participate in osmoregulation. In detailed 67 68 reviews, Péqueux (1995) and Henry (2012) assessed the specialized functions of gills and gill parts in various crustaceans and showed that both cuticle permeability and the 69

membrane characteristics of the salt-transporting gill epithelial cells were critical to
osmoregulation [1, 3]. Some bio-molecules in the gill epithelial cells facilitate salt
absorption and may also inhibit of water inflow, possibly compensating for passive
salt loss and water gain; these biomolecules include Na⁺/K⁺-ATPase, K⁺ channels, Cl⁻
channels, carbonic anhydrase, aquaporins (AQPs), and various exchangers (Na⁺/NH₄⁺,
Na⁺/H⁺, and Cl⁻/HCO₃⁻) [3, 5, 13-16].

76 The mechanisms underlying low-salinity adaptations in crustaceans have been investigated with respect to salt absorption [1, 3]. However, although the 77 78 water-permeability of epithelial cells is known to change rapidly based on the properties of some AQPs [17-18], the regulation of water inflow in invertebrates 79 (particularly crustaceans) by AQPs remains unclear [16, 19]. During osmoregulation, 80 81 it is also important to determine how energy is distributed in response to low salinity; various invertebrates have been shown to consume more energy during 82 osmoregulation [1, 3]. 83

84 The euryhaline Chinese shrimp (*Fenneropenaeus chinensis*), which has an isosmotic point of 25 ppt, is naturally distributed primarily in the Chinese Yellow Sea, 85 the Bohai Sea, and along the western coast of the Korean Peninsula [20-21]. It is an 86 important commercial shrimp along the coasts of China and Korea [20]. As these 87 88 shrimp are cultured in much lower salinity (under the isosmotic point), they must manage or tolerate substantial changes in water osmolality. In crustaceans similarly 89 exposed to salinity stress (e.g., the swimming crab, Portunus trituberculatus; the 90 Chinese crab, Eriocheir sinensis; and the Pacific white shrimp, Penaeus vannamei), 91

several genes in pathways potentially associated with adaptations to low salinity have 92 been reported [22-25]. Reports showed that F. chinensis has a poorer ability to 93 maintain a stable hemolymph osmolarity (reflecting the weaker low-salinity 94 adaptation), compared to P. vannamei [21, 26]. The poorer ability limits the farming 95 development of F. chinensis. However, to date, studies on low-salinity adaptation in F. 96 chinensis are still lacking on molecules. Recently Li (2019b) reported on F. chinensis 97 that the group after exposure at low salinity (10 ppt) showed significantly elevated 98 citrate synthase (CS) and cytochrome C oxidase (COX) activities in its gill when 99 100 compared with the group subjected to 20 ppt salinity condition [21]. For this species, these proteins have also included Na⁺/K⁺-ATPase, phenoloxidase (PO), heat shock 101 proteins (HSPs), ion-transport enzymes [27-29]. 102

103 In addition, here, we hypothesize that several other proteins, including channel transporters, AQPs, and proteins associated with energy consumption, are involved in 104 low-salinity resistance in F. chinensis. To test this hypothesis, we aimed to use 105 106 transcriptome analysis to identify and annotate the genes differentially expressed in F. chinensis exposed to low-salinity conditions, and to explore the molecular pathways 107 associated with osmoregulation or energy consumption that were overrepresented in 108 these genes. Our results will clarify the mechanisms underlying low-salinity tolerance 109 in F. chinensis and further help to explain the adaptation ability during 110 osmoregulation in this species. 111

Materials and Methods

113 Sample collection and treatment

Live shrimp were obtained from a local market near their farm of origin (in 114 Lianyungang, N 34°48'52.47", E 119°12'19.08"). Shrimp were transported to our 115 laboratory at Lianyungang Normal College. Before experimentation, all shrimp were 116 acclimated to a salinity of 20 ppt at 25°C for 5 days. This salinity was similar to the 117 natural isosmotic point of this species (25 ppt) [20, 30], as well as to the salinity at the 118 shrimp farm (23 ppt). At late stage of the acclimation period, survival rates were 119 consistently high. We then randomly divided the shrimp into two groups (n = 20 per 120 group) by two 22 L (liters) of transparent plastic tanks: the low-salinity group (LS) 121 122 was exposed to low salinity levels (5 ppt) for 24 h, while the control group remained at 20 ppt salinity as salinities ranging from 20 to 32 ppt are considered optimal 123 survival rates [20]. F. chinensis in this salinity range (20 to 32 ppt) should suffer less 124 125 salinity stress. In the study, salinity was maintained using sea salt and pure water and measured by a portable salinity meter (Arcevoos[®] ST6). In each tank, 15 L of water 126 was used and one-third of it was replaced every 12 hours (7:00-19:00). At the end of 127 the 24 h experimental period, the gills of three randomly selected surviving 128 individuals per group (mean body length: 9.13 ± 0.47 cm; mean body weight $5.13 \pm$ 129 0.65 g) were harvested and stored at -70°C for transcriptome and real-time 130 quantitative PCR (RT-qPCR) analysis. 131

132

RNA isolation, library construction, and sequencing

Total RNA was extracted using TRIzol reagent (Invitrogen Corp., USA). RNA concentration was measured using a NanoDrop 2000 spectrophotometer (Thermo Scientific, USA), and RNA integrity was assessed using 1.5% agarose gel

136	electrophoresis. Magnetic oligo (dT) beads were used to isolate mRNA from total
137	RNA. The mRNA was then fragmented into fragments approximately 200 bp long
138	using fragmentation buffer (Tris-acetate, KOAc, and MgOAc) at 94°C for 35 min.
139	The fragmented mRNA was used to construct the cDNA libraries. At least 5 μl of
140	mRNA solution (≥ 200 ng/µl) was used to construct each library. Sequencing libraries
141	for each sample were generated using the TruSeq RNA Sample Prep Kit (Illumina,
142	USA). Libraries were paired-end sequenced using a HiSeq X Ten platform (Illumina,
143	USA). The read length was 200bp.

144 **Transcriptome assembly and unigene annotation**

Raw sequence data were processed using FastqStat.jar V1.0 [31], with default 145 parameters. We then used Cutadapt v1.16 (http://cutadapt.readthedocs.io; [32]) with 146 147 parameters -q 20 -m 20 to clean the raw sequence data by deleting adapter sequences, deleting poly-N sequences, trimming low-quality sequence ends (<Q20), deleting 148 sequences with N ratios >10%, and removing reads less than 25 bp long. We used 149 Trinity (http://trinityrnaseq.github.io; [33]) to assemble the clean reads. Subsequently, 150 paired-end reads were used to fill the gaps when sequence scaffolds could not be 151 extended on either end. These sequences were defined as transcripts and were 152 subsequently assembled into unigenes based on clustering patterns using Corset [24, 153 34]. 154

The identified unigenes were annotated against six databases: NCBI nonredundant protein sequences (NR), Protein Families (PFAM), Search Tool for the Retrieval of Interacting Genes (STRING), KEGG (Kyoto Encyclopedia of Genes and

Genomes) Ortholog (KO), Gene Ontology (GO), and SWISS-PROT. We searched the
unigenes against these databases using BlastX v2.2.25 [35] with a cutoff E-value of
10⁻⁵. Functional unigenes were classified based on GO terms using Blast2GO
(http://www.blast2go.com/b2ghome) [36].

Identification and enrichment of differentially expressed unigenes (DEGs)

We used Kallisto v0.43.1 (http://kallisto.com) to evaluate the expression levels of the unigenes based on transcripts per kilobase million (TPM) values; higher TPM values reflect higher levels of unigene expression [37]. We used edgeR v3.24 to identify unigenes where $|\log 2$ fold-change (FC)| was >1 and the false discovery rate (FDR) was <0.05 [38-39]; these unigenes were considered DEGs. We then identified the KEGG pathways significantly enriched in the DEGs (P < 0.05) using hypergeometric test [40].

171 Verification of DEGs using RT-qPCR

172 We selected four RNA-Seq DEGs (three upregulated and one downregulated) for 173 RT-qPCR validation, which belong to the significant pathways (Glycine, serine and threonine metabolism, Mineral absorption and Glycolysis/Gluconeogenesis). We used 174 the β -actin gene as the internal reference gene, against which to normalize the 175 expression levels of the target genes. Gene-specific primers were designed based on 176 sequences derived from the transcriptome assembly and annotation using Primer 177 Premier 5.0 [41] (Table 1). Each RT-qPCR (25 μ l) contained 12.5 μ l of 2 × SYBR 178 qPCR Mix, 1 µl each of forward and reverse primers, 1 µl of cDNA, and 10.5 µl of 179

180	RNase-free H_2O . RT-qPCRs were performed on an Applied Biosystems 7500
181	Real-time PCR system (Applied Biosystems, Thermo Fisher Scientific, Waltham,
182	MA, USA), with the following cycling conditions: an initial denaturation step of 3
183	min at 95°C; 40 cycles of 15 s at 94°C, 15 s at 55°C, and 25 s at 72°C; and a standard
184	dissociation cycle. Three technical replicates were performed per gene, and the 2-
185	$\triangle \triangle CT$ method [42] was used to calculate relative expression levels. We considered
186	genes differentially expressed between groups if log2FC was >1 and the FDR was
187	<0.05.

188

189 190

Table 1. Primers used for real-time quantitative PCR.

Gene	Forward primer sequence	Reverse primer sequence
name	(5'-3')	(5'-3')
P. actin	GCGAGAAATCGTGCGT	AGGGTGCGAGGGCAGT
p-actin	GAC	GAT
ATP1A	AGAAGGCCGATATTGG TG	CAGGGATGTTAGATGTC AGG
CLCN2	AACAACACCCATTGCC TCAC	TCCACCAACTCCCAGAC G
IIV	GCAGACGCAGTTGACG	ACTCCTTGGCGAGACCT
НΚ	AT	Т
abil	ACAAATCTGCCCTGAA	CCTTGAACTCTGCGTCT
giya	TC	С

191

192 **Results**

193 Sequence quality and de novo assembly

After filtering the raw reads, we obtained 45,297,936 clean reads for group LS and 44,685,728 for the control group. Base call accuracy was acceptable based on Q20 values, which correspond to a 99% accuracy rate for each nucleotide base (i.e., A,

197	C, T and G) in a sequence ([43]: 97.64% and 97.16% of the bases in the LS and
198	control groups, respectively, had quality scores >20 (Q20) (Table 2). De novo
199	assembly yielded 211,955 transcripts and 159,130 unigenes. The average lengths of
200	the transcripts and unigenes were 1,028.64 bp and 662.82 bp, respectively. The N50
201	values (length of the smallest transcript/unigene in the set that contains the fewest
202	(largest) transcripts/unigenes whose combined length represents at least 50% of the
203	assembly) [44] for the transcripts and unigenes were 2,503 bp and 1,004 bp,
204	respectively (Table 3). The raw data have been uploaded to SRA database and the
205	BioProject ID is PRJNA669213.

Table 2. Clean-read statistics.

Group	Total reads	Total bases	Error%	Q20%	GC%
Low-salinity (LS)	45,297,936	6,725,104,745	0.03	97.14	45.99
Control	44,685,728	6,634,797,034	0.03	97.16	47.40

Table 3.	De novo	assembly	statistics.
1 4010 0		assembly	Statistics.

Туре	Unigenes	Transcripts
Total sequence num	159,130	211,955
Total sequence base	105,474,756	218,024,931
Percent GC	42.42	43.36
Largest (bp)	38,633	38,633
Smallest (bp)	201	186
Average (bp)	662.82	1028.64
N50	1,004	2,503
N90	269	341

212 Annotation and classification of the transcriptome

213	We were able to annotate only a small fraction of the 159,130 unigenes against
214	the six databases (10.50% on average). The greatest proportion of unigenes (13.99%;
215	22,256 unigenes) was annotated against the NR database, followed by the STRING
216	and SWISS-PROT databases (10.89% and 10.94%, respectively; Table 4). GO
217	analysis of the annotated unigenes showed that, in the biological process category, the
218	three terms annotated in the most unigenes were macromolecule metabolic process
219	(10,578 unigenes), organonitrogen compound metabolic process (10,297 unigenes),
220	and regulation of cellular process (9,871 unigenes); in the cellular components
221	category, the three terms annotated in the most unigenes were intracellular (14,934
222	unigenes), intracellular part (14,883 unigenes), and cytoplasm (14,078 unigenes); and
223	in the molecular function category, the three terms annotated in the most unigenes
224	were cation binding (5,800 unigenes), nucleic acid binding (5,521 unigenes), and
225	anion binding (5,322 unigenes) (Fig 1).

226

227 228 Table 4. Annotation statistics of 159 130 Unigenes on F. chinensis.

Databasa	No. of	% of
Database	unigenes	total
GO	16,578	10.42%
KEGG	13,975	8.78%
NR	22,256	13.99%
PFAM	13,360	8.40%
STRING	17,325	10.89%
SWISS-PROT	16,700	10.49%
Average	16.699	10.50%

229

230 DEG identification and enrichment

We identified 3,658 unigenes as DEGs (i.e., |log2FC |>1 and FDR < 0.05). Of

these, 1,755 were downregulated in the LS group as compared to the control, and 232 1,903 were upregulated (Fig 2). We identified 38 KEGG pathways as significantly 233 234 enriched in the 3,658 DEGs (P < 0.05, Fig 3). Of these, 13 were metabolic pathways, including tryptophan metabolism, lysine degradation, drug metabolism-cytochrome 235 P450, and tyrosine metabolism; 13 were organismal systems pathways, including 236 salivary secretion, insulin secretion, proximal tubule bicarbonate reclamation, and the 237 TOLL and IMD signaling pathways; and two were cellular process pathways, namely 238 phagosome and regulation of the actin cytoskeleton (Fig 3). The remaining 10 239 240 pathways were associated with environmental information processing, drug development, and human diseases (Fig 3). Across all 38 pathways, three were 241 potentially related to osmoregulation: mineral absorption (ko04978), four DEGs; 242 243 vasopressin-regulated water reabsorption (KEGG: ko04962), three DEGs; and ribosome (ko03010), 29 DEGs (Table 5). The DEGs in these pathways included 244 ATP1A (Sodium/potassium-transporting ATPase subunit alpha), CLCN2 (Chloride 245 246 channel 2), VAMP2 (Vesicle-associated membrane protein 2), and AOP4 (Aquaporin-4), HMOX2 (Heme oxygenase 2), SLC40A1/ FPN1 (Solute carrier family 247 40 (iron-regulated transporter), member 1), RAB5 (Ras-related protein Rab-5), etc. 248 (Table 5). 249

250

251 Table 5. Unigenes in osmoregulation-related pathways differentially expressed in

response to low salinity. OS: Organismal Systems; GIP: Genetic Information Processing

Gene Symbol	Up/Down-regulati	Description	KEGG ID
	on (log2FC), LS		

	vs Control					
Mineral absorption-KEGG pathway (OS)						
ATP1A	Upregulated	Sodium/potassium-transporting ATPase	ATPase			
	(2.49)	subunit alpha				
CLCN2	Upregulated	Chloride channel 2	ClC-2			
	(1.87)					
HMOX2	Upregulated	Heme oxygenase 2	HMOX			
	(1.10)					
SLC40A1/FPN1	Downregulated	Solute carrier family 40 (iron-regulated	FPN1			
	(-1.18)	transporter), member 1				
Vasopressin-regulated wate	er reabsorption- KEGC	G pathway (OS)				
AQP4	Downregulated	Aquaporin-4;	AQP4			
	(-1.14)					
RAB5	Downregulated	Ras-related protein Rab-5	Rab5			
	(-1.26)					
VAMP2	Downregulated	Vesicle-associated membrane protein 2	VAMP2			
	(-1.20)					
Ribosome- KEGG pathway	(GIP)					
RP-L40e	Upregulated	Large subunit ribosomal protein L40e	L40e			
	(1.30)					
<i>RP-L36</i>	Upregulated	Large subunit ribosomal protein L36e	L36e			
	(1.22)					
RP-L7	Upregulated	Large subunit ribosomal protein L7/L12	L7/L12			
~	(1.21)		~ ~			
RP-S3e	Downregulated	Small subunit ribosomal protein S3e	S3e			
	(-1.10)					
RP-S4e	Downregulated	Small subunit ribosomal protein S4e	S4e			
	(-1.03)		60			
RP-59	Downregulated	Small subunit ribosomal protein 89	89			
	(-1.57)		014			
<i>KP-S14e</i>	Downregulated	Small subunit ribosomal protein S14e	S14e			
	(-1.06)		015-			
KP-SIJe	(1.12)	Small subunit ribosomal protein \$15e	515e			
DD \$154a	(-1.13)	Small subunit ribosomal protain \$154 a	Q15 A o			
KF-SIJAe	(1 20)	Sman subunit ribosomai protein STSAe	SIJAe			
DD \$10a	(-1.20)	Small subunit ribosomal protain \$10a	S10 e			
M -519e	(1.07)	Sinan subunit noosoniai protein 319e	5196			
RP_S21_a	(-1.07)	Small subunit ribosomal protein \$21e	\$210			
M -521e	(-1.38)	Sinan subunit noosoniai protein 521e	5210			
RP-S23p	Downregulated	Small subunit ribosomal protein \$23e	\$23e			
14 5250	(-1 06)	Shan Subunt Hobsoniai protein 525e	5250			
RP-S27e	Downregulated	Small subunit ribosomal protein S27e	S27e			
	(-1.24)	·····	-			

RP-S29e	Downregulated (-1.06)	Small subunit ribosomal protein S29e	S29e
RP-L5e	Downregulated (-1.10)	Large subunit ribosomal protein L5e	L5e
RP-L7e	Downregulated (-1.05)	Large subunit ribosomal protein L7e	L7e
RP-L9e	Downregulated (-1.27)	Large subunit ribosomal protein L9e	L9e
RP-L10Ae	Downregulated (-1.15)	Large subunit ribosomal protein L10Ae	L10Ae
RP-L13e	Downregulated (-2.71)	Large subunit ribosomal protein L13e	L13e
RP-L14e	Downregulated (-1.00)	Large subunit ribosomal protein L14e	L14e
RP-L17e	Downregulated (-1.10)	Large subunit ribosomal protein L17e	L17e
RP-L18e	Downregulated (-1.40)	Large subunit ribosomal protein L18e	L18e
RP-L21e	Downregulated (-1.07)	Large subunit ribosomal protein L21e	L21e
RP-L23e	Downregulated (-2.59)	Large subunit ribosomal protein L23e	L23e
RP-L28e	Downregulated (-1.07)	Large subunit ribosomal protein L28e	L28e
RP-L29e	Downregulated (-1.02)	Large subunit ribosomal protein L29e	L29e
RP-L31e	Downregulated (-1.11)	Large subunit ribosomal protein L31e	L31e
RP-L35Ae	Downregulated (-1.12)	Large subunit ribosomal protein L35Ae	L35Ae
RP-L35e	Downregulated (-1.10)	Large subunit ribosomal protein L35e	L35e
RP-L37Ae	Downregulated (-1.92)	Large subunit ribosomal protein L37Ae	L37Ae
RP-L37e	Downregulated (-1.04)	Large subunit ribosomal protein L37e	L37e

253

254 **RT-qPCR verification**

255 We used RT-qPCR to verify four DEGs: two DEGs from osmoregulation-related 256 pathways (*ATP1A* and *CLCN2*; Table 5) and two random DEGs in other pathways

(*HK* and *glyA*). Consistent with the RNA-seq results, RT-qPCR identified *ATP1A*, *CLCN2*, and *HK* (hexokinase) as significantly upregulated in the LS group as compared to the control (|log2FC| > 1 and FDR < 0.05; Fig 4). Although *glyA* (glycine hydroxymethyltransferase) was not significantly downregulated between the LS and control groups in the qRT-PCR analysis, this gene had similar patterns of expression in both the qRT-PCR and the RNA-Seq analyses (Fig 4).

263 **Discussion**

264 Assembly quality and GO classification

265 The F. chinensis transcriptome assembled in this study had an N50 of 1,004 bp, which was similar to, but slightly better than, those previously obtained for P. 266 vannamei (448 bp; [23]) and Oratosquilla oratoria (798 bp; [45]). This indicated that 267 268 our assembly was of acceptable quality. Consistent with previous studies of osmoregulation ([23-24]), the unigenes identified in this study were primarily 269 associated with cation binding, macromolecule metabolic process, and organonitrogen 270 271 compound metabolic process. This suggested that genes with functions in these categories are potentially important to adaptation to low-salinity environments. The 272 result also partially supports the finding from Yuan (2017), who showed that 15.02 % 273 and 16.24 % of positively selected genes in seawater and freshwater shrimps, 274 respectively, enriched in the functions of cation binding; 22.71% and 18.80% 275 positively selected genes in seawater and freshwater shrimps, respectively, enriched in 276 277 the functions of cellular macromolecule metabolic process [46]. In their study, however, no specific and osmoregulation-related data for F. chinensis is available, 278

thus genes used in this study are not identical compared to that in their study. Whether unigens enriched in other GO terms mentioned by this study (Fig 1) have positive selection sites to adapt the low-salinity environments, which is another question to answer in future.

DEGs and pathways adapted to low salinity

Minerals not only serve as nutrients, but also are essential components of 284 osmoregulation for crustaceans [3]. In the LS group, three genes in the mineral 285 absorption pathway (ko04978) were differentially expressed as compared to the 286 control: three were upregulated (ATP1A, CLCN2, and HMOX2), and one was 287 downregulated (SLC40A1/FPN1; Table 5). Of these, ATP1A, CLCN2, and 288 SLC40A/FPN1 encode channel transporter proteins, while HMOX2 encodes an 289 290 intracellular protein. Our identification of these DEGs in this important pathway suggested that they may play a role in osmoregulation in response to low-salinity 291 exposure. 292

ATP1A is involved in encoding a Na⁺/K⁺-ATPase that controls the movements of 293 the Na^+ and K^+ ions between the hemolymph and the intracellular fluid [1]. The 294 upregulation of Na⁺/K⁺-ATPase on the basolateral membrane causes more Na⁺ ions to 295 be transported out than K⁺ ions taken in [47]. The upregulation of the chloride channel 296 protein (encoded by CLCN2) has a similar effect on osmoregulation, increasing the 297 amount of Cl⁻ leaving the cell and entering the hemolymph space. This process thus 298 facilitates the rapid adaptation of F. chinensis to low-salinity environments by 299 increasing salt concentration in hemolymph [1]. Indeed, previous studies have shown 300

that F. chinensis hemolymph is hyperosmotic to the external medium at low salinities 301 (e.g., 5 ppt) [10, 30]. On this point, the results support the opinion reviewed by 302 Péqueux (1995) and Henry (2012) that osmoregulators in low-salinity environment 303 (below 26 ppt) will turn on mechanisms of anisomotic extracelluar regulation to 304 stabilize hemolymph osmotic and ionic concentrations [1, 3]. Although the control 305 group (20 ppt) is already in low-salinity environment according to reference salinity 306 of 26 ppt, the result has implied that the salinity difference between 5 ppt and 20 ppt 307 is too huge enough to make their gene expression difference, as well as the 308 309 hemolymph osmolality [21].

We also observed that two genes associated with iron levels were differentially 310 expressed in the LS group as compared to the control: *HMOX2*, which encodes heme 311 312 oxygenase 2, was upregulated, and SLC40A1/FPN1, which encodes an iron-regulated transporter, was downregulated. The differential expression of these genes may lead 313 to the increased production of ferrous irons and reduced iron outflow (ko04978) in the 314 315 LS group. A previous study had showed that the decreased iron concentration in blood of Cacinus maenas was associated with their adaptation to osmotic stress [48]. The 316 downregulation of the iron-regulated transporter (DIRT) in this study has provided a 317 new interpretation that how the iron concentration in blood was decreased at the 318 molecular level. Besides the crustaceans, the DIRT even occurred in fish species like 319 steelhead trout (Oncorhynchus mykiss). However, fewer reports clearly showed the 320 function of the iron in osmoregulation [48-49]. Decreasing blood iron concentration in 321 crustaceans under salinity stress was interpreted by the sortation of iron from blood to 322

other tissues [48]. Moreover, extreme salinity stress will generates an increase in the ROS (reactive oxygen species) which is harmful to crustaceans [3]. Iron in tissues plays a role in oxidative metabolism by the form of a key competent in cytochromes and enzymes [50]. Thus, taken together, it may be the way for *F. chinensis* in low salinity in response to salinity stress that the iron in gill cells is prone to be used for synthesis of bio-molecules (cytochromes and enzymes) involved in oxidative metabolism.

In addition regulating salt and mineral levels, crustaceans maintain an 330 331 approximately constant osmotic concentration of extracellular fluid (hemolymph) regardless of the salinity of the surrounding medium, by regulating water flow in and 332 out of the hemolymph [1]. Here, three unigenes in the vasopressin-regulated water 333 reabsorption pathway (ko04962; AQP4, VAMP2, and RAB5) were downregulated in 334 the LS group. We expect that this downregulation will reduce water inflow to the 335 hemolymph, helping to maintain a constant osmotic concentration. In particular, 336 because the primary function of AQP4 is to transport water across the plasma 337 membrane into hemolymph [19, 51], thus, the downregulation of the AOP4 gene will 338 facilitate reduction of hemodilution. Similarly, the downregulation of VAMP2 will 339 strongly inhibit AQP2 fusion at the apical membrane, which has been shown to 340 decrease water flow into the hemolymph in vertebrates [18, 52]. Finally, the 341 downregulation of *RAB5* may also inhibit the fusion of AQP2 at the endosomal apical 342 membrane; RAB5 is also one of the components implicated in early endosome fusion 343 [53], particularly, which is predicted to be involved in the regulation of AQP2 344

trafficking to and from the plasma membrane [54]. Thus, this study reports, for the
first time, that the genes (*AQP4*, *VAMP2*, and *RAB5*) are associated with the reduction
of water-permeability in *F. chinensis* in response to low-salinity environments.

In the study, we expected that the allostatic load on F. chinensis would increase 348 when salinity decreased from 20 ppt to 5 ppt. All forms of allostasis require energy, as 349 allostatic load increases, the amount of energy available for other biological functions 350 decreases [6, 21, 23, 25, 55]. We found that most of the DEGs in the ribosome 351 pathway (ko03010) were downregulated in the LS group as compared to the control 352 353 (Table 5). Ribosome is the location of polypeptide synthesis. Downregulation of the structural macromolecular components in ribosome could decrease polypeptide 354 synthesis. The previous study showed proteins L4, L22, L39e, L19, L23, L24, L29 355 356 and L31e are important to polypeptide synthesis [56]. Downregulation of these genes in this study implies the synthesis of proteins will be affected in F. chinensis exposed 357 to low salinities. Notably, it seems that proteins not involved in low-salinity resistance 358 more likely decrease, concomitant with the diversion of energy resources to 359 osmoregulation. Similarly, P. vannamei subjected to chronic low-salinity stress 360 upregulated the expression of AMP-activated protein kinase (AMPK) to maintain 361 energy balance by increasing catabolism to generate ATP and decreasing anabolism to 362 conserve ATP [57]. However, AMPK was not significantly differentially expressed in 363 F. chinensis. This suggested that, unlike P. vannamei (another common shrimp 364 species cultured in China), F. chinensis may not have a strong ability to adapt to 365 low-salinity conditions in maintaining energy balance. Otherwise, another evidence 366

implying the weaker adaption of *F. chinensis* to low salinity compared to *P. vannamei*can be found in studies from Tang (2016) and Li (2019b) [21, 26]. Hemolymph
osmolality of *F. chinensis* was significantly reduced when salinity decreased from 15
ppt to 10 ppt [21], while that of *P. vannamei* did not significantly reduced even when
environmental salinity decreased from 12 ppt to 0 ppt [26]. Thus, successfully
low-salinity aquaculture of *F. chinensis* deserves more attention.

In this study, we used two different salinities to identify DEGs potentially associated with the response of *F. chinensis* to low salinity. Because we only compared two salinities, our results do not reflect the adaptation process. To better understand the mechanisms associated with gradual or continuous changes in salinity, we aim to investigate the responses of this species to additional salinities in future studies, using both RNA-Seq and RT-qPCR.

In summary, our results indicated that four unigenes in the mineral absorption 379 pathway (ATP1A, CLCN2, HMOX2, and SLC40A1/FPN1), as well as three unigenes 380 in the vasopressin-regulated water reabsorption pathway (AOP4, VAMP2, and RAB5), 381 were differentially expressed in F. chinensis in response to low-salinity exposure. 382 These pathways, in conjunction with the ribosome pathway, may be important for 383 osmoregulation in F. chinensis under low-salinity conditions. Although the associated 384 mechanisms require further investigation, our results help to clarify the molecular 385 responses of F. chinensis to low-salinity environments. This study suggests that F. 386 chinensis could be an evolutionary model of weake osmoregulator, combining with 387 patterns of hemolymph osmoregulation (include the strong osmoregulator, weak 388

osmoregulator and osmoconformer, etc.) in aquatic crustaceans viewed by Péqueux

390 (1995) [1].

J31 Data Availability Statement

The raw data have been uploaded to SRA database (PRJNA669213).

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408 **Competing Interests**

409 The authors have declared that no competing interests exist.

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412 **References**

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79-90.

Fig 1. Gene Ontology (GO) analysis of the unigenes in the *Fenneropenaeus chinensis* genome, showing the GO terms most overrepresented in the GO categories (A) biological processes, (B) cellular components, and (C) molecular function. The colors in each pie chart correspond to GO terms, and the size of each slice represents the proportion of unigenes associated with each GO term.

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630 Fig 2. Unigene expression patterns. The horizontal and vertical axes present the expression levels of unigenes in the two groups (TPM, transcripts per kilobase 631 632 million, values); each value was logarithmically transformed. DEGs, i.e., unigenes with expression fold changes $|\log 2FC| > 1$ and FDR < 0.05, are marked with red and 633 blue. Red dots represent unigenes that were significantly upregulated under low 634 salinity, blue dots represent those that were significantly downregulated under low 635 salinity, and black dots represent genes that are not DEGs. The greater the deviation 636 of a dot from the diagonal, the greater the difference in the unigene expression 637 between the two groups. Dots near 0 represent unigenes with low expression. 638

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Fig 3. KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways significantly
enriched in the DEGs. Each bar represents a pathway, and the height of bar reflects
the enrichment ratio (equal to Sample Number / Background Number). *: FDR <
0.05, **: FDR < 0.01.

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Fig 4. RT-qPCR verification of four representative genes (*ATP1A*, *CLCN2*, *HK*, and *glyA*) identified as differentially expressed between low-salinity and control groups of *Fenneropenaeus chinensis*. *: FDR < 0.05. Each bar with standard error represents three replicates.



(B)





Expression level of unigenes in the low-salinity group



