

***Litopenaeus vannamei* stylicins are constitutively produced by hemocytes and intestinal cells and are differentially modulated upon infections**

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Abstract :

Stylicins are anionic antimicrobial host defense peptides (AAMPs) composed of a proline-rich N-terminal region and a C-terminal portion containing 13 conserved cysteine residues. Here, we have increased our knowledge about these unexplored crustacean AAMPs by the characterization of novel stylicin members in the most cultivated penaeid shrimp, *Litopenaeus vannamei*. We showed that the *L. vannamei* stylicin family is composed of two members (*Lvan-Stylicin1* and *Lvan-Stylicin2*) encoded by different loci which vary in gene copy number. Unlike the other three gene-encoded antimicrobial peptide families from penaeid shrimp, the expression of *Lvan-Stylicins* is not restricted to hemocytes. Indeed, they are also produced by the columnar epithelial cells lining the midgut and its anterior caecum. Interestingly, *Lvan-Stylicins* are simultaneously transcribed at different transcriptional levels in a single shrimp and are differentially modulated in hemocytes after infections. While the expression of both genes showed to be responsive to damage-associated molecular patterns, only *Lvan-Stylicin2* was induced after a *Vibrio* infection. Besides, *Lvan-Stylicins* also showed a distinct pattern of gene expression in the three portions of the midgut (anterior, middle and posterior) and during shrimp development. We provide here the first evidence of the diversity of the stylicin antimicrobial peptide family in terms of sequence and gene expression distribution and regulation.

Highlights

- Stylicins from *L. vannamei* are composed of two members, *Lvan-Stylicin1* and -2.
- *Lvan-Stylicins* are encoded by separate genes and transcribed at different levels.
- *Lvan-Stylicins* are expressed in hemocytes and in midgut epithelial cells.
- Expression of *Lvan-Stylicins* is differentially modulated in response to infections.

Keywords : Crustacean, Penaeid shrimp, Invertebrate immunity, Antimicrobial peptide, Host defense peptide, Molecular diversity

45 **1. Introduction**

46 Antimicrobial host defense peptides (AMPs) are important components of the
47 innate immune system of both vertebrates and invertebrates. They are usually described
48 as gene-encoded peptides (less than 10 kDa) with cationic and amphipathic properties
49 which selectively target the negatively charged membranes of microbes [1]. In addition
50 to those classical cationic antimicrobial peptides (also known as CAMPs), the current
51 classification of AMPs also includes polypeptides/proteins larger than 10 kDa, AMPs
52 generated by the processing of precursor molecules and anionic peptides [2]. Anionic
53 antimicrobial peptides (AAMPs) comprise a non-phylogenetic group of either gene-
54 encoded or non-ribosomally synthesized molecules with a high proportion of anionic
55 amino acid residues (aspartate and glutamate). AAMPs are widely distributed in living
56 organisms and play an important role in host defense against bacteria, fungi and viruses
57 [3]. Like their cationic counterparts, AAMPs are multifunctional molecules engaged in
58 different biological and immunological processes beyond antimicrobial functions [3].

59 Shrimp farming is an important economic activity for many developing countries
60 in Asia and Latin America, which has been repeatedly threatened by infections caused
61 by viruses and pathogenic bacteria from the genus *Vibrio*. Consequently, infectious
62 disease outbreaks are clearly a major concern in aquaculture that has encouraged
63 extensive research efforts. The scientific findings in the last decade have provided
64 valuable information on the role of AMPs in shrimp defenses. More than natural
65 antibiotics, shrimp AMPs are also involved in the control of the natural microbiota,
66 wound healing, bacterial clearance and other immunomodulatory functions [4]. To date,
67 four gene-encoded AMP families have been identified in the hemocytes of penaeid
68 shrimp: penaeidins, crustins, anti-lipopolysaccharide factors (ALFs) and stylicins [4,5].

69 Stylicins were initially identified as transcripts associated to shrimp survival to
70 pathogenic *Vibrio* infections [6]. Characterization of stylicins revealed that they are
71 anionic peptides of 8.9 kDa composed of an N-terminal proline-rich region followed by
72 a C-terminal region containing 13 cysteine residues [7]. In *Litopenaeus stylirostris*
73 shrimp, stylicins form a diverse AAMP family composed of two members, named *Lsty-*
74 *Stylicin1* and *Lsty-Stylicin2*. Although the antimicrobial activity of the recombinant
75 r*Lsty-Stylicin1* has been shown to be restricted to filamentous fungi, the r*Lsty-Stylicin1*
76 displayed a strong lipopolysaccharide (LPS)-binding activity and the ability to
77 agglutinate Gram-negative bacteria [7]. In the kuruma prawn *Marsupenaeus japonicus*,
78 the gene expression of its single stylicin (*Mjap-Stylicin*) showed to be modulated in
79 gills and hepatopancreas in response to the White spot syndrome virus (WSSV) [8].
80 Curiously, apart from these two reports [7,8], no other stylicin members have been
81 characterized thus far.

82 In order to fill this research gap, novel members of the stylicin AMP family were
83 identified and characterized in the most important cultured penaeid species, *Litopenaeus*
84 *vannamei* (*Lvan-Stylicin1* and *Lvan-Stylicin2*). We showed that *Lvan-Stylicins* (also
85 known as *Vibrio* penaeicida-induced cysteine and proline-rich peptides or *LvVICPs* [9])
86 are highly anionic peptides encoded by two distinct genomic loci that follow different
87 patterns of gene regulation during shrimp development and after microbial infections.
88 Interestingly, while both genes responded to danger/damage-associated molecular
89 patterns (shrimp muscle tissues), only *Lvan-Stylicin2* showed to be up-regulated in
90 circulating hemocytes in response to a *Vibrio* infection. Moreover, by combining
91 immunohistochemistry and whole-mount immunofluorescence assays, we showed that
92 *Lvan-Stylicins* are also constitutively produced by the midgut columnar epithelial cells.

93 To our knowledge, this is the first evidence for the expression of a shrimp gene-encoded
94 AMP in other tissues than the immune cells from the hemolymph.

95

96 **2. Materials and Methods**

97 **2.1. Animals, immune challenge and tissue collection**

98 Juvenile Pacific white shrimp (*Litopenaeus vannamei*) (10 ± 2 g) were obtained
99 from the Laboratory of Marine Shrimp of the Federal University of Santa Catarina
100 (Southern Brazil). After an acclimation period of seven days, animals (n=5) were
101 stimulated by the injection of 5×10^7 colony-forming units (CFU)/animal of heat killed
102 (70°C for 20 min) *Vibrio harveyi* ATCC 14126 in 100 μL sterile seawater (SSW).
103 Naïve (unchallenged) animals (n=5) were used as control. At 48 h post-stimulation,
104 hemolymph was withdrawn into modified Alsever solution (MAS: 27 mM sodium
105 citrate, 336 mM NaCl, 115 mM glucose, 9 mM EDTA, pH 7.0) and hemocytes were
106 separated from plasma by centrifugation ($800 \times g$ for 10 min at 4°C). Shrimp were
107 subsequently anesthetized (ice bath for 10 min) and sacrificed for the collection of the
108 following tissues: gills, muscle, nerve cord, hepatopancreas, foregut, midgut and
109 hindgut. Tissue samples were washed in Tris-saline solution (10 mM Tris, 330 mM
110 NaCl, pH 7.4), homogenized in TRIzol reagent (Thermo Scientific) and immediately
111 processed for total RNA isolation and tissue distribution analysis. Pleopods from naïve
112 shrimp were collected and processed for genomic DNA (gDNA) extraction.

113

114 **2.2. Experimental infections**

115 Two unrelated shrimp pathogens were chosen for experimental infections, the
116 Gram-negative *Vibrio harveyi* and the White spot syndrome virus (WSSV). For the

117 bacterial infection, 6×10^7 CFU/animal of live *V. harveyi* ATCC 14126 in 100 µL SSW
118 (median lethal dose within 2 days, LD50/2) or 100 µL SSW (injury control) were
119 injected. For the viral infection, shrimp were injected with 100 µL of a WSSV inoculum
120 containing 3×10^2 viral particles (median lethal dose within 15 days, LD50/15). The
121 WSSV inoculum was prepared from muscle tissues of WSSV-infected shrimp as
122 previously described [10]. Control animals for the viral infection were injected with 100
123 µL of a muscle tissue homogenate prepared from WSSV-free shrimp. At 48 h post-
124 infections, circulating hemocytes and midguts were collected, pooled (3 pools of 5
125 animals per condition) and processed for total RNA extraction and quantitative PCR
126 analysis of gene expression. Unchallenged animals (naïve shrimp at time 0 h) were used
127 as control for all experiments.

128 In a second experiment, individual shrimp (n=3) were challenged by the oral
129 administration of 7.5×10^5 CFU/animal of live *V. harveyi* ATCC 14126 in 50 µL SSW.
130 Shrimp (n=3) that received SSW served as controls. The bacterial *per os* challenge was
131 performed as previously described [11]. No mortalities were recorded during the course
132 of the experiment. At 21 h post-challenge, midguts from both *Vibrio*-challenged and
133 unchallenged shrimp were collected and flushed with cold Tris-saline solution and then
134 cut into three equal portions (anterior, middle and posterior). Each individual sample
135 was homogenized in TRIzol reagent (Thermo Scientific) and processed for total RNA
136 extraction and quantitative PCR analysis of gene expression.

137

138 **2.3. Genomic DNA and total RNA extraction and cDNA synthesis**

139 For gDNA extraction, individual pleopods were homogenized and incubated at
140 55°C for 1 h in 500 µL of lysis buffer (100 mM Tris-HCl pH 8.5, 100 mM NaCl, 50

141 mM EDTA pH 8.0, 1% SDS, 0.25 µg/µL proteinase K). After addition of 3 M
142 potassium acetate (1:2; v:v), samples were incubated at 4°C for 30 min and centrifuged
143 at 14,000 ×g for 10 min. Following precipitation with isopropanol, gDNA samples were
144 washed in 70% ethanol and treated with 50 µg/mL RNase A (Fermentas) at 37°C for 30
145 min. Quantification and quality of gDNA samples were assessed by spectrophotometry
146 and 0.8% agarose gel electrophoresis, respectively.

147 Total RNA was extracted using TRIzol reagent (Thermo Scientific) according to
148 the manufacturer's instructions. RNA samples were treated with DNase I (Thermo
149 Scientific) at 37°C for 15 min and precipitated with 0.3 M sodium acetate (pH 5.2) and
150 isopropanol (1:1; v:v). RNA amount and quality were assessed by spectrophotometric
151 analysis and the integrity of total RNA was analyzed by 0.8% agarose gel
152 electrophoresis. Following heat denaturation (70°C for 5 min), reverse transcription was
153 performed using 1 µg of purified total RNA with 50 ng/µL oligo(dT)₁₂₋₁₈ in a 20-µL
154 reaction volume containing the RevertAid Reverse Transcriptase (Thermo Scientific),
155 according to the manufacturer's instructions.

156

157 **2.4. Molecular cloning**

158 PCR amplifications for molecular cloning were conducted using primers based on
159 the nucleotide sequence of two stylicin homologues (contigs: DN31608_c0_g1_i1 and
160 DN31608_c0_g1_i2) identified in midgut transcriptomes of *L. vannamei* (unpublished
161 data). PCR reactions were carried out in a 15-µL reaction volume containing 50-100 ng
162 of gDNA, 2 mM MgCl₂, 0.4 mM dNTP Mix, 0.4 µM of each primer (Table 1) and 1 U
163 Taq DNA Polymerase (Sinapse). PCR conditions were as follows: 1 cycle of
164 denaturation at 95°C for 10 min followed by 35 cycles of 95°C for 45 s, 55°C for 45 s

165 and 72°C for 2 min, and a final extension step of 72°C for 10 min. The amplification
166 products were analyzed by electrophoresis (1.5% agarose gel) and cloned into a
167 pCR2.1-TOPO vector (Thermo Scientific). The positive clones were identified by
168 colony PCR and plasmid sequencing.

169

170 **2.5. Sequence data analysis and phylogeny**

171 Stylicin sequences from the penaeid species *L. vannamei* (contigs:
172 DN31608_c0_g1_i1 and DN31608_c0_g1_i2), *L. stylirostris* (*Lsty*-Stylicin1:
173 EU177435; *Lsty*-Stylicin2: EU177437) and *M. japonicus* (*Mjap*-Stylicin: KR063277)
174 were used for the search of homologous sequences in the following publicly accessible
175 databases: Expressed Sequence Tags (EST), Transcriptome Shotgun Assembly (TSA)
176 and Whole-Genome Shotgun Contigs (WGS). Only full-length coding sequences were
177 included. Homology searches were performed using tBLASTX at the NCBI web servers
178 (<http://www.ncbi.nlm.nih.gov/BLAST>). All nucleotide sequences were manually
179 inspected and translated using the ExPASy Translate Tool
180 (<http://web.expasy.org/translate/>). Prediction of signal peptide was performed with the
181 SignalP 4.1 program (<http://www.cbs.dtu.dk/services/SignalP/>) and the theoretical
182 isoelectric point (pI) and molecular weight (MW) of the mature peptides were predicted
183 using the ExPASy ProtParam Tool (<http://web.expasy.org/protparam/>). Phylogenetic
184 analysis based on both nucleotide and predicted amino acid sequences were conducted
185 in MEGA X [12] using the Maximum Likelihood method. Bootstrap sampling was
186 reiterated 1,000 times using a 50% bootstrap cutoff.

187

188 **2.6. Semiquantitative RT-PCR analysis**

189 The extraction of total RNA and cDNA synthesis were performed using the
190 method described above. PCR reactions were carried out in a 15- μ L reaction volume
191 containing 1 μ L of cDNA, 2 mM MgCl₂, 0.4 mM dNTP Mix, 0.4 μ M of each primer
192 (Table 1) and 1 U Taq DNA Polymerase (Sinapse). PCR conditions were as follows: 1
193 cycle of denaturation at 95°C for 5 min followed by 30 cycles of 95°C for 30 s, 57°C for
194 30 s and 72°C for 30 s, and a final extension step of 72°C for 10 min. PCR products
195 were analyzed by electrophoresis (1.5% agarose gel) and stained by ethidium bromide.
196 The expression of the *LvActin* gene (PCR conditions: 40 cycles of 95°C for 45 s, 50°C
197 for 45 s and 72°C for 1 min) was used as endogenous control to normalize the RT-PCR
198 data for comparison.

199

200 **2.7. Immunodetection of stylicins in shrimp tissues**

201 Whole juvenile shrimp (n=3) were fixed in Davidson's fixative solution (22%
202 formalin, 31.5% ethanol and 11.5% glacial acetic acid) for 24 h at room temperature,
203 embedded in paraffin and cut into 5 μ m thick sections. Histological sections of shrimp
204 tissues were deparaffinized and hydrated through xylene-ethanol-water series, and
205 washed in Tris-buffered saline (TBS: 50 mM Tris-HCl, 200 mM NaCl, pH 7.2). Then,
206 sections were permeabilized (1×TBS, 0.1% Triton X-100) for 30 min and blocked in
207 TBS-T solution (1×TBS, 1% BSA, 0.05% Tween 20) for 2 h followed by 16 h
208 incubation at 4°C with mouse anti-r*Lsty*-Stylicin1 polyclonal antibodies (2.3 μ g/mL)
209 [7]. After three washes in 1×TBS+0.05% Tween 20 buffer, sections were incubated for
210 3 h at room temperature with alkaline phosphatase-labeled rabbit anti-mouse IgG
211 (1:1000) (Thermo Scientific), followed by a 1 h incubation at room temperature in the
212 dark in a solution of 100 mM Tris-HCl, 100 mM NaCl, 50 mM MgCl₂ (pH 9.3)

213 containing 0.175 mg/mL 5-bromo-4-chloro-3-indolyl phosphate (BCIP) (Sigma), 0.375
214 mg/mL nitro blue tetrazolium (NBT) (Sigma) and 0.24 mg/mL levamisole (Sigma).
215 Negative controls consisted in replacing anti-r*Lsty*-Stylicin1 antibodies with pre-
216 immune mouse serum or TBS-T solution.

217 Whole-mount immunofluorescence assays were conducted in midgut samples
218 from juvenile shrimp (n=3). Midguts were harvested by dissection, washed in ice-cold
219 Tris-saline solution and immediately fixed in 4% paraformaldehyde. Just after removal
220 of the intestinal content, midguts were longitudinally opened, washed in phosphate-
221 buffered saline (PBS: 137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 2 mM KH₂PO₄,
222 pH 7.2) and blocked in PBS-T solution (1×PBS, 1% BSA, 0.1% Triton X-100) for 3 h
223 followed by 16 h incubation at 4°C with mouse anti-r*Lsty*-Stylicin1 polyclonal
224 antibodies (2.3 µg/mL) [7]. Midguts were then washed 3 times with PBS-T solution for
225 20 min and incubated for 4 h at room temperature with 4',6-diamidino-2-phenylindole
226 (DAPI) and FITC-conjugated anti-mouse secondary antibodies diluted at 1:500
227 (Thermo Scientific). Negative controls consisted in replacing the primary antibodies
228 with pre-immune mouse serum or PBS-T solution. The experiments were repeated three
229 times, and representative images were taken by confocal microscope (Leica DMI6000 B
230 Microscope).

231

232 **2.8. Fluorescence-based reverse transcription real-time quantitative PCR (RT-
233 qPCR)**

234 RT-qPCR reactions were performed in a final volume of 15 µL containing 0.2 µM
235 of each primer (Table 1), 7.5 µL of Maxima SYBR Green/ROX qPCR Master Mix
236 (Thermo Scientific) and 1 µL of cDNA. The RT-qPCR program was 95°C for 10 min,

237 followed by 40 cycles of 95°C for 15 s and 60°C for 60 s. Melt curve analysis (60-95°C
238 at a temperature transition rate of 0.05°C/s) was performed to evaluate primer
239 specificity. Primer pair efficiencies (E) were calculated from 2-fold dilution series of
240 pooled cDNA for each primer pair. Primer pair efficiencies were calculated from the
241 given slopes in the StepOne software v2.3, according to the equation: $E = 10^{(-1/\text{slope})}$.

242 The eukaryotic translation elongation factor 1-alpha (*LvEF1α*) and the ribosomal
243 proteins *LvL40*, *LvRpS3A* and *LvRpS6* (Table 1) were used as reference genes of
244 expression data in circulating hemocytes. In midgut, *LvEF1α*, *LvL40* and *LvActin* were
245 used as reference genes for data normalization [13]. The relative expression levels of
246 *Lvan-Stylicins* were calibrated with the expression profile of circulating hemocytes or
247 midgut from naïve (unchallenged) shrimp, according to the $2^{-\Delta\Delta C_q}$ method [14].
248 Differences in gene expression were considered statistically significant at $P < 0.05$
249 (cutoff of 1.5-fold change in expression levels) using one-way ANOVA and Tukey's
250 multiple comparison test.

251 The transcript abundance of *Lvan-Stylicins* in three midgut portions (anterior,
252 middle and posterior) from *Vibrio*-challenged and naïve (unchallenged) shrimp was
253 quantified by RT-qPCR and normalized with the gene expression of *LvEF1α*, *LvRpS3A*,
254 *LvRpS6* and *LvActin* (Table 1). The relative expression levels were calibrated with the
255 gene expression of each midgut portion from unchallenged shrimp and differences were
256 considered statistically significant at $P < 0.05$ (cutoff of 1.5-fold change in expression
257 levels) using Student's t-test.

258

259 **2.9. Gene copy number estimation and basal mRNA levels**

260 The number of *Lvan-STY* gene copies in *L. vannamei* genome was estimated in
261 five individual shrimp by absolute quantification through the qPCR technique. qPCR
262 reactions were performed using specific primers (Table 1) and 80 ng of gDNA as
263 template. The absolute quantification of the target genes (*Lvan-STY1* and *Lvan-STY2*)
264 was calculated using a standard curve derived from 10-fold dilution series of plasmids
265 containing the DNA target sequences (10^7 to 10^3 plasmids/ μ L; $R^2= 0.998$ and 0.999 for
266 *Lvan-STY1* and *Lvan-STY2*, respectively). The transcript abundance of *Lvan-Stylicin1*
267 and *Lvan-Stylicin2* in the circulating hemocytes of the same five shrimp individuals was
268 assessed by absolute quantification using 1 ng of reverse-transcribed total RNA. The
269 absolute mRNA quantification was performed using the same method described for the
270 gene copy number estimation.

271 The comparison of the transcript abundance of *Lvan-Stylicins* between hemocytes
272 and midgut was assessed by RT-qPCR (3 pools of 4 animals per tissue) and normalized
273 with the gene expression of *LvEF1 α* , *LvRpS3A* and *LvRpS6* (Table 1), according to the
274 $2^{-\Delta\Delta C_q}$ method [14]. Differences in gene expression between tissues were considered
275 statistically significant at $P < 0.05$ (cutoff of 1.5-fold change in expression levels) using
276 Student's t-test.

277

278 **2.10. Quantitative gene expression analysis during shrimp development**

279 Three biological replicates of different development stages of *L. vannamei* were
280 collected: fertilized eggs at 0-4 h (EI) and at 7-11 h post-spawning (EII), nauplius I and
281 V (NI and NV), protozoaea I and III (ZI and ZIII), mysis I and III (MI and MIII) and
282 postlarvae aged of 2, 9 and 17 days (PL2, PL9 and PL17). Transcript levels of *L.*
283 *vannamei* stylicins during shrimp development were quantified by RT-qPCR and

normalized with the gene expression of *LvRpS6* and *LvActin* (Table 1), as previously described [15]. Hemocyte samples from juveniles (3 pools of 5 animals) were used as control for calibrating gene expression data. Genes were considered as “not expressed” in a specific development stage when PCR amplification yielded no product (no dissociation curves) whereas RT-qPCR reactions showing Cq values higher than the limit of quantification (but that generated an expected dissociation curve profile) were considered as “unquantifiable”. Statistical significance was considered at $P < 0.05$ by one-way ANOVA followed by Tukey’s multiple comparison test.

292

293 3. Results

294 3.1. *L. vannamei* stylicins comprise a diverse AAMP family

295 The transcriptomic analysis of the *L. vannamei* midgut (unpublished) has revealed
296 the presence of two nucleotide sequences (contigs: DN31608_c0_g1_i1 and
297 DN31608_c0_g1_i2) homologous to the stylicin antimicrobial peptides from the blue
298 shrimp *L. stylirostris* (*Lsty*-Stylicin1 and *Lsty*-Stylicin2). Both sequences correspond to
299 full-length transcripts that encode for precursors composed of a signal peptide followed
300 by a mature peptide containing 13 cysteine residues. The full-length stylicin sequences
301 from *L. vannamei* were cloned by PCR amplification from gDNA samples and re-
302 sequenced for confirmation. The nucleotide sequence corresponding to the contig
303 DN31608_c0_g1_i2 shared 86% identity with *Lsty*-Stylicin1 (GenBank: EU177435)
304 whereas the contig DN31608_c0_g1_i1 shared 93% identity with *Lsty*-Stylicin2
305 (GenBank: EU177437). Thus, those sequences from *L. vannamei* were designated as
306 *Lvan*-Stylicin1 (GenBank: MH108957) and *Lvan*-Stylicin2 (GenBank: MH108958),
307 respectively (Fig. 1A).

308 The deduced amino acid sequences of *Lvan*-Stylicins start with a predicted 22-
309 residue signal peptide followed by an anionic mature peptide of 82 amino acid residues
310 (Fig. 1A; Table S1). Although *Lvan*-Stylicins contain no obvious protein domains, both
311 mature peptides are characterized by the presence of a proline-rich N-terminal region
312 and a C-terminal region holding the 13 conserved cysteine residues found in *L.*
313 *stylirostris* stylicins and in the *Mjap*-Stylicin from *M. japonicus*. *Lvan*-Stylicins were
314 quite similar to each other and to stylicins from *L. stylirostris* (>80% identity), but less
315 than 65% amino acid identity was observed between *Lvan*-Stylicins and *Mjap*-Stylicin.

316 *In silico* analysis of *L. vannamei* non-annotated databases publicly available on
317 GenBank led to the identification of different isoforms for each *L. vannamei* stylicin:
318 five for *Lvan*-Stylicin1 (GenBank: FE179060, FE156583, FE124653, FE125173 and
319 GETD01027084) and three for *Lvan*-Stylicin2 (GenBank: GETD01027083,
320 GETZ01053100 and GFRP01011277). Those sequences were identified in EST and
321 TSA libraries from multiple shrimp tissues, such as hemocytes, lymphoid organ, nerve
322 cord and hepatopancreas. Additionally, we included in this analysis the *Vibrio*
323 *penaeicida*-induced cysteine and proline-rich peptides (*LvVICP1* and *LvVICP2*),
324 stylicin homologues identified in *L. vannamei* by Wang and colleagues [9]. While *Lvan*-
325 Stylicin2 sequences differed only by synonymous and non-synonymous substitutions,
326 *Lvan*-Stylicin1 sequences showed two distinct lengths: 82 or 83 amino acid residues
327 (Fig. 1B; Table S1). Besides from the synonymous and non-synonymous substitutions,
328 *Lvan*-Stylicin1 peptides can be distinguished from each other by the presence/absence
329 of a tyrosine (Y) or histidine (H) residue at the position 51 of the precursor peptide (Fig.
330 1B). The 82-residue *Lvan*-Stylicin1 peptides showed a molecular weight of 8.83 kDa
331 and a calculated pI of 4.98 while the 83-residue *Lvan*-Stylicin1 peptides showed a

332 molecular weight ranging from 8.97 to 9.02 kDa and a calculated pI of 4.98 or 5.19
333 (Table S1). Comparatively, the mature *Lvan*-Stylicin2 peptides had a molecular weight
334 of about 9 kDa and a calculated pI ranging from 4.47 to 4.69 (Table S1).

335

336 **3.2. *L. vannamei* stylicins are encoded by distinct genomic loci**

337 The genomic organization of *L. vannamei* stylicin genes (*Lvan-STY1* and *Lvan-*
338 *STY2*) was investigated in two individual shrimp using PCR-based and cloning
339 strategies. Sanger DNA sequencing results showed that *Lvan*-Stylicins are encoded by
340 distinct genomic sequences, but share a similar structural gene organization (Fig. 2A).
341 *Lvan-STY1* and *Lvan-STY2* genes are composed by two exons interrupted by a single
342 intron with a length of 217 bp and 205 bp, respectively (Fig. 2A). Both genomic DNA
343 sequences followed the canonical GT/AG splicing recognition rule at the exon/intron
344 boundaries (Fig. S1). In both genes, the first exon covers the 5'-untranslated region
345 (UTR), the signal peptide and the first three residues of the mature peptide while the
346 second exon encodes the remainder of the mature peptide sequence and the 3'-UTR.

347 Interestingly, the *Lvan-STY1* genes from the two sampled shrimp were identical to
348 each other while the nucleotide sequence of their *Lvan-STY2* genes differed in ten
349 nucleotides: one in the first exon, five in the intron and four in the second exon (Fig.
350 S1). The five nucleotide substitutions found in the coding sequence resulted in the
351 change of two amino acid residues. The obtained genomic sequences were deposited in
352 GenBank under the accession numbers MH108959 to MH108962.

353 Then, we asked whether *Lvan-STY1* and *Lvan-STY2* genes have the same number
354 of copies in *L. vannamei* genome. The relative gene copy number ratio of *Lvan-*
355 *STY2/Lvan-STY1* was estimated in five individual animals by quantitative PCR. The

356 number of *Lvan-STY2* gene copies was 2.37 ± 0.15 -fold higher than the number of copies
357 of the *Lvan-STY1* gene (Fig. 2B). Finally, a phylogenetic analysis showed that the *Lvan-*
358 *STY* genes were placed in the same clade with the stylicin genes from *L. stylirostris*
359 (*Lsty-STY1*: EU177436 and *Lsty-STY2*: EU177437) (Fig. 2C). In this clade, *Lvan-STY1*
360 and *Lsty-STY1* clustered together in a single group distinct to the *STY2* genes (Fig. 2C).
361 Stylicin genes from *P. monodon* (*Pmon-STY*: NIUS012084699) and *M. japonicus*
362 (*Mjap-STY*: NIUR011088360) clustered in separate groups.

363

364 **3.3. Stylicins cluster into three distinct phylogenetic groups**

365 *In silico* mining of publicly accessible databases (EST, TSA and WGS) resulted in
366 the identification of novel members of the Stylicin family in different penaeid species:
367 *Fenneropenaeus penicillatus* (*Fpen-Stylicin*: GFRT01005742), *M. japonicus* (*Mjap-*
368 *Stylicin*: NIUR011088360) and *P. monodon* (*Pmon-Stylicin*: JZ892895, DW678047,
369 DW678039, DT366712, DW042940, GW996588, GEEP01015864, GEME01013089
370 and NIUS012084699). From our *in silico* analysis, stylicin sequences were only
371 identified in penaeid shrimp species. All obtained sequences hold the 13 conserved
372 cysteine residues at the C-terminal region (Fig. 3A).

373 Phylogenetic analysis revealed that the Stylicin family is a monophyletic group
374 that evolved from a common ancestor gene. Phylogenetic trees constructed with
375 nucleotide and predicted amino acid sequences shared similar topological structures. As
376 shown in Fig. 3B, stylicins clustered in three main clades. The first clade included only
377 stylicin sequences from penaeid species from the genus *Litopenaeus*. In this clade,
378 Stylicin1 and Stylicin2 were split into two distinct groups (Fig. 3B). Finally, while

379 *Pmon*-Stylicins and *Fpen*-Stylicin clustered in a second phylogenetic group, *Mjap*-
380 Stylicins formed a separate clade from all stylicin sequences (Fig. 3B).
381

382 **3.4. Stylicins are constitutively produced by hemocytes and intestinal cells**

383 The gene expression distribution of *L. vannamei* stylicins was first evaluated in
384 eight different tissues of shrimp stimulated or not by the injection of heat-killed *V.*
385 *harveyi*. Results from the semiquantitative RT-PCR analysis evidenced the presence of
386 *Lvan*-Stylicin1 transcripts in circulating hemocytes, foregut, midgut and gills (Fig. 4). In
387 *Vibrio*-stimulated animals, *Lvan*-Stylicin1 gene expression was also detected in hindgut
388 and nerve cord. Comparatively, transcripts of *Lvan*-Stylicin2 were detected in
389 circulating hemocytes, foregut, midgut, hindgut, gills and nerve cord of both stimulated
390 and non-stimulated animals (Fig. 4). For both *Lvan*-Stylicins, no signals were observed
391 in muscle and hepatopancreas (Fig. 4).

392 To characterize the peptide localization of *Lvan*-Stylicins, immunohistochemistry
393 analysis was subsequently performed on sections of different shrimp tissues using
394 polyclonal antibodies raised against the r*Lsty*-Stylicin1 from *L. stylrostris* [7]. Due to
395 the high degree of sequence conservation, anti-*Lsty*-Stylicin1 antibodies probably
396 recognized both *Lvan*-Stylicins. Stylicin immunoreactivity was found in individual cells
397 heterogeneously distributed across the shrimp tissues. In the cephalothorax, positive
398 immunoreactivity was especially pronounced in some cells present in the connective
399 tissue of the anterior midgut caecum (Fig. 5A-B). Based on morphological features,
400 those stylicin-positive cells are likely tissue-infiltrating hemocytes.

401 Besides those cells, stylicin immunoreactivity was also observed in the apical
402 region of the columnar epithelial cells lining the midgut and its anterior caecum (Fig.

403 5A-B). Whole-mount immunofluorescence assays were further performed to confirm
404 the presence of stylicin peptides in those epithelial cells. Confocal images clearly
405 evidenced the presence of stylicin-containing granules located at the apical region of the
406 midgut columnar epithelial cells (Fig. 5C-E). No signals were observed in other cell
407 types of the shrimp body, thus the results of the semiquantitative RT-PCR analysis are
408 probably the consequence of the infiltration of stylicin-expressing hemocytes in shrimp
409 tissues. Altogether, results from both immunohistochemistry and whole-mount
410 immunofluorescence assays showed that *Lvan*-Stylicins are constitutively produced by
411 the hemocytes and by the columnar epithelial cells of the midgut.

412

413 **3.5. *Lvan-STY1* and *Lvan-STY2* genes are simultaneously transcribed in a single**
414 **shrimp at different basal levels**

415 Since *Lvan*-Stylicins are produced by both hemocytes and midgut cells, we
416 focused on determining the main site of stylicin expression in penaeid shrimp. Results
417 showed that the expression of *Lvan*-Stylicin1 and *Lvan*-Stylicin2 was, respectively,
418 52.5-fold and 62.2-fold higher in circulating hemocytes than in the midgut (Fig. 6A).
419 Next, the basal mRNA expression levels of the *Lvan-STY1* and *Lvan-STY2* genes were
420 analyzed in the circulating hemocytes of five individual shrimp by absolute
421 quantification. Interestingly, both *Lvan-STY* genes showed to be constitutively and
422 simultaneously transcribed in an individual shrimp, but at different transcriptional
423 levels. The basal expression of *Lvan-STY2* was 3.69-fold higher than *Lvan-STY1* (Fig.
424 6B). Besides, the basal mRNA levels of each gene showed to be also variable among
425 the individuals (Fig. 6B). For the *Lvan-STY1* gene, differences in gene expression

426 reached up to 3.78-fold whereas variations up to 2.36-fold were found in *Lvan-STY2*
427 gene expression.

428

429 **3.6. The gene expression of *L. vannamei* stylicins is differentially modulated in**
430 **response to infections**

431 The gene expression profile of *Lvan*-Stylicins was further quantified by
432 fluorescence-based quantitative PCR (RT-qPCR) in shrimp hemocytes and midgut at 48
433 h after infections with two unrelated pathogens, the Gram-negative *V. harveyi* and the
434 WSSV. This time point was chosen on the basis of previous studies from our group
435 [10,13]. Interestingly, while the expression of *Lvan*-Stylicin2 was induced in circulating
436 hemocytes in response to the *Vibrio* infection (2.4-fold change), the expression of *Lvan*-
437 Stylicin1 was not modulated (Fig. 7). No increase in *Lvan*-Stylicin2 expression was
438 observed following the injection of sterile seawater (aseptic injury control). Besides,
439 both genes were not modulated after the viral infection. In contrast, the expression of
440 both genes was up-regulated in circulating hemocytes (2.5-fold change for *Lvan*-
441 Stylicin1 and 2.9-fold change for *Lvan*-Stylicin2) after the injection of a muscle tissue
442 homogenate prepared from WSSV-free shrimp (Fig. 7). In the midgut, the expression of
443 *Lvan*-Stylicins was not regulated by the bacterial or by the viral infection (Fig. 7).

444 The lack of gene expression response in the midgut encouraged us to conduct an
445 alternative experimental infection method, mimicking a more natural route of bacterial
446 infection (*per os* challenge). Variations in gene expression of *Lvan*-Stylicins were
447 assessed by RT-qPCR in three midgut portions (anterior, middle and posterior). The
448 expression of both *Lvan*-Stylicins was up-regulated (2.89-fold change for *Lvan*-
449 Stylicin1 and 4.31-fold change for *Lvan*-Stylicin2) only in posterior portion of the

450 midgut (Fig. 8). By contrast, no increase in *Lvan-Stylicin* gene expression was observed
451 in the two first midgut portions (anterior and middle) in response to the oral *Vibrio*
452 challenge (Fig. 8).

453

454 **3.7. *L. vannamei* stylicins show a different pattern of gene expression during**
455 **shrimp development**

456 We finally investigated the presence and the levels of stylicin transcripts in twelve
457 developmental stages of *L. vannamei*, from fertilized eggs to larval (nauplius, protozoaea
458 and mysis) and postlarval stages, and also in circulating hemocytes from juveniles.
459 Transcript levels of *Lvan-Stylicin1* were detected at very low levels in late protozoaea
460 stages (ZIII), but its expression was only quantified from mysis III (MIII). Then, *Lvan-*
461 *Stylicin1* expression increased gradually in the following developmental stages (Fig. 9).
462 On the other hand, *Lvan-Stylicin2* transcripts were found to be present early in shrimp
463 development (fertilized eggs at 7-11 h post-spawning). However, *Lvan-Stylicin2*
464 expression was only quantified from the protozoaea III (ZIII) stage (Fig. 9). For both
465 genes, the highest mRNA abundance was quantified in hemocytes from juvenile shrimp
466 (Fig. 9).

467

468 **4. Discussion**

469 We showed here that *L. vannamei* stylicins comprise a diverse family of anionic
470 antimicrobial peptides (AAMPs) whose genes are differentially regulated in hemocytes
471 and midgut cells in response to infections. From the four gene-encoded AMPs described
472 in penaeid shrimp, only stylicins have not been fully characterized in the Pacific white
473 shrimp (*L. vannamei*) and this is the first study exploring the diversity of the stylicin

family in terms of sequence and gene expression distribution and regulation. By taking advantage of RNA-Seq technology, we have identified two stylicin homologues (*Lvan-Stylicin1* and *Lvan-Stylicin2*) in midgut transcriptomes of *L. vannamei* showing high similarities to stylicins from the blue shrimp *L. stylirostris* [7]. From our *in silico* analysis, stylicins form a diverse gene family in shrimp species of the genus *Litopenaeus*, but not in other taxa of the family Penaeidae, such as *Fenneropenaeus*, *Penaeus* and *Marsupenaeus*. Essentially, our results revealed that all known *Litopenaeus* gene-encoded AMPs are present as diverse multigene families composed of different members. For instance, while *Litopenaeus* penaeidins are composed of three members (*Litvan PEN1/2*, *Litvan PEN3* and *Litvan PEN4*), at least four and seven members were identified in the crustin (Crustin *Lv*, Crustin-like *Lv*, *LvSWD* and *LvSPLI*) and in the ALF families (*Litvan ALF-A* to *-G*), respectively [4,5,16]. However, unlike other gene-encoded AMPs from marine invertebrates, stylicins are exclusively composed of anionic peptides. Interestingly, the spectrum of activity of these anionic antimicrobials is restricted to filamentous fungi [7], even if their gene expression has shown to be associated to shrimp survival to pathogenic *Vibrio* infections [6]. Likewise, anionic peptides derived from the C-terminus of the shrimp respiratory protein hemocyanin are also specific against fungi [17]. These hemocyanin-derived peptides, named PvHCT, are able to bind and permeabilize fungal membranes [18]. To date, the mechanism of action of stylicins and of other AAMPs is completely unknown. Actually, in comparison to CAMPs, few scientific groups have attempted to study the subject of AAMPs [3] and more functional studies are needed to achieve a more in-depth understanding of these unconventional AMPs.

497 We showed here that *Lvan-Stylicin1* and *Lvan-Stylicin2* are encoded by different
498 genomic loci. Nonetheless, the presence of distinct stylicin genes was only observed in
499 *Litopenaeus* species. Thus, it is likely that *STY1* and *STY2* are paralogous genes that
500 arose from a single gene duplication event before the speciation of the genus
501 *Litopenaeus*. Besides, although both *Lvan-STY* genes share a similar structural gene
502 organization, their copy numbers in *L. vannamei* genome showed to be variable,
503 suggesting that they have followed independent duplication events after *STY1-STY2*
504 divergence. Apart from the two main stylicins members (*Lvan-Stylicin1* and *Lvan-*
505 *Stylicin2*), we have identified *Lvan-Stylicin1* sequences with distinct lengths (82 and 83
506 amino acids in length). No evidences for the presence of alternative splicing were found
507 in the *Lvan-STY1* gene, and thus it is most likely that different alleles may occur for *L.*
508 *vannamei* stylicin genes. Taken together, our results suggest that *L. vannamei* stylicins
509 belong to a diverse multigenic and multiallelic family of AAMPs. Moreover, *Lvan-*
510 *STY1* and *Lvan-STY2* genes showed to be simultaneously transcribed in a single shrimp.
511 The different penaeidin and ALF members are also simultaneously expressed in an
512 individual shrimp [16,19], and it would be of great interest to colocalize the four gene-
513 encoded AMP families in shrimp hemocytes and examine whether they function
514 synergistically to enhance their antimicrobial spectrum of activity. Synergic activities
515 have already been observed for AMPs from both vertebrate [20] and invertebrate
516 species [21,22], but unfortunately, no studies of this type have been addressed in
517 crustaceans.

518 One of the most important findings of this study is that the expression *Lvan-*
519 *Stylicins* is not limited to the immune cells (hemocytes) as observed for the other three
520 shrimp gene-encoded AMPs [4]. Both immunostaining and gene expression analysis

revealed that *Lvan*-Stylicins are constitutively produced by the midgut columnar epithelial cells and that their expression is induced in response to *Vibrio* infections. Notably, the expression of *Lvan*-Stylicin was pronounced in the anterior midgut caecum, an intestinal region primarily involved in the production, secretion and activation of digestive enzymes [23]. Thus, it is plausible to hypothesize that *Lvan*-Stylicins are secreted into the midgut lumen. This finding brings new insights into the role of crustacean AMPs in the control of the gut microbiota and in shrimp intestinal defenses. Indeed, the shrimp midgut lacks the cuticular lining found in the other portions of the intestine (foregut and hindgut), representing a potential route of entry for many pathogens into the hemocel. In a previous study, we showed that gut is an important source for the expression of immune-related genes in penaeid shrimp [13], thus the presence of stylicins in the midgut cells suggests the participation of these antimicrobial effectors in the first intestinal line of defense. Interestingly, the expression of some ALF members showed to be involved in the maintenance of the microbiota residing in the shrimp hemolymph [24]. Taken all together, our results highlight the importance of stylicins in both hemolymph-based and gut-based immunities.

Another relevant conclusion that can be drawn from this study is that *Lvan*-STY genes are differentially regulated after infections. Such diversity in gene expression regulation was previously reported for the different members of the *L. vannamei* ALF family in response to fungal infections [25]. Interestingly, while some AMPs such as *Lvan*-Stylicin1, penaeidins [26], *Litvan* ALF-A [25] and Type II crustins [27] are not-regulated in response to infections, the expression of other *L. vannamei* AMPs (*Lvan*-Stylicin2 and other ALF members) is induced in immune cells to improve host antimicrobial responses against pathogens. As elegantly shown by Wang *et al.* [9], the

545 gene expression of most shrimp AMPs, including penaeidins, Type II crustins and
546 stylicins (*LvVICPs*), is controlled by the IMD/NF- κ B pathway, an evolutionarily
547 conserved signaling cascade involved in the regulation of the antimicrobial responses of
548 arthropods. On the other hand, the expression of almost all shrimp AMPs, including
549 *Lvan*-Stylicins, can be drastically affected in circulating hemocytes during lethal
550 infections [10]. Unlike the *Mjap*-Stylicin from the kuruma prawn *M. japonicus* [8],
551 *Lvan*-Stylicins were not modulated neither in circulating hemocytes nor in midgut by
552 the WSSV. This result could be probably due to the time course response of *Lvan*-
553 Stylicins in those shrimp tissues. On the other hand, both *Lvan*-STY genes have been
554 shown to be responsive to a tissue homogenate prepared from shrimp muscle (injury
555 control for the WSSV infection). This finding strongly suggests that *Lvan*-Stylicins are
556 induced in response to danger/damage-associated molecular patterns (DAMPs) and that
557 these AAMPs could be involved in early inflammation and in wound healing processes
558 as proposed for penaeidins [26].

559 *L. vannamei* stylicin genes showed to be differentially regulated not only in
560 response to infections, but also during shrimp development. Interestingly, *Lvan*-
561 Stylicin2 and the *M. japonicus* stylicin showed a very similar pattern of gene expression
562 during shrimp development [8]. Comparatively, *Lvan*-Stylicin1 expression was quite
563 similar to that observed for other *L. vannamei* AMPs, such as *Litvan* PEN1/2, *Litvan*
564 PEN4 and *Litvan* ALF-D [15]. Apart from that, we cannot discard the possibility of
565 these changes in gene expression could be the result of the differences in the mRNA
566 basal levels observed between the *Lvan*-STY genes. Indeed, the detection of stylicin
567 transcripts, as well as of other gene-encoded AMPs, highlights the importance of these
568 antimicrobial effectors during shrimp development.

569

570 **5. Conclusions**

571 In conclusion, we showed that the stylicin family from shrimp species of the
572 genus *Litopenaeus* is composed of two members encoded by distinct genomic loci that
573 exhibit different patterns of gene expression distribution and regulation. According to
574 the best of our knowledge, this is the first evidence for the expression of a shrimp gene-
575 encoded AMP in other tissues than the hemocytes. Even though the expression of
576 stylicins has been shown to be a marker of shrimp survival to pathogenic *Vibrio*
577 infections, the role of these effectors in shrimp immune defense is still largely unknown.
578 The application of RNAi-based methods could make significant contributions to
579 understanding the significance of the molecular and transcriptional diversity of
580 *Litopenaeus* stylicins in host-microbe interactions.

581

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594

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691

692 **Appendix A. Supplementary data**

693 **Supplemental Table S1.** Biochemical properties of stylicin antimicrobial peptides from
694 penaeid shrimp.

695 **Supplemental Figure S1.** Multiple nucleotide alignment of *Lvan-STY1* and *Lvan-STY2*
696 genes from two single shrimp: individual #1 (*Lvan-STY1*: MH108959; *Lvan-STY2*:
697 MH108960) and individual #2 (*Lvan-STY1*: MH108961; *Lvan-STY2*: MH108962). Exon
698 sequences are in bold. Nucleotide substitutions and the canonical GT/AG splice site
699 junctions are shadowed with black and gray backgrounds, respectively.

700

701 **Tables**702 **Table 1.** Nucleotide sequences of primers used in this study.

Gene	Forward primer (5'-3')	Reverse primer (5'-3')	Amplicon
Primers for molecular cloning and sequencing			
<i>Lvan-STY1</i>	CTGGACGCATCCCTGCTG	TGCGCCTTCGTTCTCTTATCC	571 bp
<i>Lvan-STY2</i>	GCTGTACTGCTCCTGTGTAG	CTTCGTTCTCGCTTCTTATCC	589 bp
Primers for tissue distribution analysis (RT-PCR)			
<i>Lvan-Stylecin1</i>	CACAAGAGTGCCCACCGTG	ACATTCCGCAGTTATGGTAGCC	125 bp
<i>Lvan-Stylecin2</i>	CACAAGAGTGCCCACCGTG	CACACAGGCTGCCGACATAA	151 bp
<i>LvActin</i>	TAATCCACATCTGCTGGAAGGTGG	TCACCAACTGGGATGACATGG	846 bp
Primers for absolute and relative quantification analyses (qPCR and RT-qPCR)			
<i>Lvan-Stylecin1</i>	CACAAGAGTGCCCACCGTG	ACATTCCGCAGTTATGGTAGCC	125 bp
<i>Lvan-Stylecin2</i>	CACAAGAGTGCCCACCGTG	CACACAGGCTGCCGACATAA	151 bp
<i>LvActin</i>	CCACGAGACCACCTACAAC	AGCGAGGGCAGTGATTTC	142 bp
<i>LvEF1α</i>	TGGCTGTGAACAAGATGGACA	TTGTAGCCCACCTTCTTGACG	103 bp
<i>LvL40</i>	GAGAATGTGAAGGCCAAGATC	TCAGAGAGAGTGCAGACCATC	104 bp
<i>LvRpS3A</i>	GGCTTGCTATGGTGTGCTCC	TCATGCTCTGGCTCGCTG	101 bp
<i>LvRpS6</i>	AGCAGATAACCCTGGTGAAG	GATGCAACCACGGACTGAC	193 bp

703

704

705 **Figure captions**

706 **Figure 1.** (A) Nucleotide and deduced amino acid sequences (one letter code) of *Lvan-*
 707 *Stylicin1* (GenBank: MH108957) and *Lvan-Stylicin2* (GenBank: MH108958). The
 708 predicted signal peptides are in bold and underlined. Asterisks (*) mark the stop codon.
 709 (B) Amino acid sequence alignments of stylicins from penaeid shrimp species from the
 710 genus *Litopenaeus* (*Lvan*: *L. vannamei* and *Lsty*: *L. stylirostris*). Identical amino acid
 711 residues are highlighted in black while specific amino acid residues found in Stylicin1
 712 and Stylicin2 peptides are highlighted in blue and yellow, respectively. Triangles (▼)
 713 indicate the 13 conserved cysteine residues. GenBank accession numbers are indicated
 714 in brackets. The sequences of the *Vibrio penaeicida*-induced cysteine and proline-rich
 715 peptides (*LvVICP1* and *LvVICP2*) were obtained from [9].

716

717 **Figure 2.** (A) Not-to-scale schematic representation of stylicin genes from *Litopenaeus*
 718 *vannamei* (*Lvan-STY1* and *Lvan-STY2*). Boxes represent the exons and the line between
 719 boxes represents the intron. Numbers indicate the length of exons and introns in base
 720 pairs. (B) Estimation of the copy number of *Lvan-STY* genes in five individual shrimp.
 721 The absolute quantification was assessed by qPCR using a standard curve derived from
 722 10-fold dilution series of plasmids containing each target gene. Results are presented as
 723 the ratio of the abundance of *Lvan-STY2* gene copies per ng of gDNA to that of *Lvan-*
 724 *STY1*. (C) Structural organization and phylogenetic relationship of *STY* genes from
 725 penaeid shrimp: *Litopenaeus vannamei* (*Lvan-STY1*: MH108959 and MH108961; *Lvan-*
 726 *STY2*: MH108960 and MH108962), *Litopenaeus stylirostris* (*Lsty-STY1*: EU177436;
 727 *Lsty-STY2*: EU177437), *Penaeus monodon* (*Pmon-STY*: NIUS012084699) and
 728 *Marsupenaeus japonicus* (*Mjap-STY*: NIUR011088360). The cladogram at the left of

729 the figure indicates the phylogenetic relationship of *STY* genes. Pink and yellow boxes
730 indicate the position of the signal peptides and the mature stylicins in the exons,
731 respectively, while the black lines indicate the introns.

732

733 **Figure 3.** (A) Amino acid sequence alignments of mature peptides of stylicins. Identical
734 amino acid residues are highlighted in black. Triangles (▼) indicate the 13 conserved
735 cysteine residues. GenBank accession numbers are indicated in brackets. (B)
736 Phylogenetic analysis of stylicins from penaeid shrimp. The tree was constructed using
737 the Maximum Likelihood method with bootstrap values calculated from 1,000 trees.
738 Sequences included in analyses were the following: *Litopenaeus vannamei* (*Lvan-*
739 *Stylicin1*: MH108957, MH108959, MH108961, FE179060, FE156583, FE124653,
740 FE125173, GETD01027084 and *LvVICP1*; *Lvan-Stylicin2*: MH108958, MH108960,
741 MH108962, GETD01027083, GETZ01053100, GFRP01011277 and *LvVICP2*),
742 *Litopenaeus stylirostris* (*Lsty-Stylicin1*: EU177435, EU177436 and GM615897; *Lsty-*
743 *Stylicin2*: EU177437), *Fenneropenaeus penicillatus* (*Fpen-Stylicin*: GFRT01005742),
744 *Penaeus monodon* (*Pmon-Stylicin*: JZ892895, DW678047, DW678039, DT366712,
745 DW042940, GW996588, GEEP01015864, GEME01013089 and NIUS012084699) and
746 *Marsupenaeus japonicus* (*Mjap-Stylicin*: KR063277 and NIUR011088360).

747

748 **Figure 4.** Gene expression distribution of *Litopenaeus vannamei* stylicins (*Lvan-*
749 *Stylicin1* and *Lvan-Stylicin2*) in different tissues from naïve (N) and *Vibrio*-stimulated
750 (S) shrimp. Gene expression analysis was performed by semiquantitative RT-PCR using
751 the β -actin gene (*LvActin*) as an endogenous expression control. The figure (not-to-
752 scale) shown at the top of the figure indicates the anatomic location of shrimp tissues:

753 foregut (FG), hepatopancreas (HP), midgut (MG), hindgut (HG), circulating hemocytes
754 (HE), muscle (ML), gills (GL) and nerve cord (NC).

755

756 **Figure 5.** (A) Immunodetection of *Lvan-Stylicin* peptides in shrimp tissues by
757 immunohistochemistry. Stylicin immune reactivity was observed in hemocytes
758 infiltrating the connective tissue (arrows) and in the columnar epithelium of the midgut.
759 The figure (not-to-scale) shown at the top panel indicates the anatomic location of
760 shrimp tissues: anterior midgut caecum (AMC), midgut (MG) and hepatopancreas (HP).
761 Scale bars = 100 µm. (B) A magnification of the anterior midgut caecum. The arrows
762 indicate stylicin-expressing hemocytes infiltrating connective tissues. Negative controls
763 consisted in replacing primary antibodies with pre-immune mouse serum (bottom
764 panel). Scale bars = 100 µm. (C) Scanning confocal microscopy images of the
765 immunodetection (whole mount immunofluorescence staining) of *Lvan-Stylicin*
766 peptides present in granules located at the apical region of the midgut columnar
767 epithelial cells. (D) Nuclei of the midgut columnar epithelial cells stained with DAPI.
768 (E) Merged images of the stylicin-containing granules (green) and the nuclei (blue) of
769 the midgut columnar epithelial cells. Scale bars = 20 µm.

770

771 **Figure 6.** (A) Quantitative comparison of the relative abundance of *Lvan-Stylicin1*
772 (white bars) and *Lvan-Stylicin2* (black bars) transcripts in circulating hemocytes (HE)
773 and midgut (MG). Results are presented as mean ± standard deviation of relative
774 expressions (three biological replicates) and statistical differences are indicated by
775 asterisks (*) (Student's t-test, $P < 0.05$). (B) Transcript abundance of *Lvan-Stylicins* in
776 five individual shrimp. The absolute quantification was assessed by qPCR using a

777 standard curve derived from 10-fold dilution series of plasmids containing each target
778 gene. Results are presented as the ratio of the abundance of *Lvan-Stylicin2* transcripts
779 per ng of total RNA to that of *Lvan-Stylicin1*. The β-actin gene (*LvActin*) was used as
780 endogenous expression control for each individual shrimp (lower panel).

781

782 **Figure 7.** Relative gene expression profile of *Lvan-Stylicin1* (white bars) and *Lvan-*
783 *Stylicin2* (black bars) in circulating hemocytes and midgut of shrimp at 48 h after
784 experimental infections with the Gram-negative *Vibrio harveyi* ATCC 14126 (6×10^7
785 CFU/animal) or the White spot syndrome virus (WSSV: 3×10^2 viral particles/animal).
786 Results are presented as mean \pm standard deviation of relative expressions (three
787 biological replicates) and statistical differences are indicated by asterisks (*) (one-way
788 ANOVA/Tukey, $P < 0.05$). W-free: tissue homogenate inoculum prepared from WSSV-
789 free shrimp.

790

791 **Figure 8.** Relative gene expression profile of *Lvan-Stylicin1* and *Lvan-Stylicin2* in three
792 portions of shrimp midgut (anterior, middle and posterior) at 21 h after the oral
793 administration of 7.5×10^5 CFU/animal of live *V. harveyi* ATCC 14126. Results are
794 presented as mean \pm standard deviation of relative expressions (three biological
795 replicates) and statistical differences are indicated by asterisks (*) (Student's t-test, $P <$
796 0.05). The figure (not-to-scale) shown at the top of the figure indicates the anatomic
797 location of the three midgut portions from naïve (white bars) and *Vibrio*-challenged
798 (black bars) shrimp.

799

800 **Figure 9.** Gene expression profile of *Litopenaeus vannamei* stylicins during shrimp
801 development. EI: fertilized eggs at 0-4 h post-spawning; EII: fertilized eggs at 7-11 h
802 post-spawning; NI: nauplius I; NV: nauplius V; ZI: protozoa I; ZIII: protozoa III; MI:
803 mysis I; MIII: mysis III; PL2: postlarva 2; PL9: postlarva 9; PL17: postlarva 17. The
804 red dotted line indicates the basal expression level in hemocytes from juvenile shrimp
805 while the solid blue underline highlights the stages at which the gene expression was
806 detected (valid dissociation curve profile) but not quantified (Cq values higher than the
807 limit of quantification). Results are presented as mean \pm standard deviation (three
808 biological replicates). Different letters indicate significant differences among the
809 development stages while asterisks (*) shows significant differences between each
810 developmental stage and hemocytes from juveniles (one-way ANOVA/Tukey, $P <$
811 0.05).

A

Lvan-Stylicin1

Lvan-Stylicin2

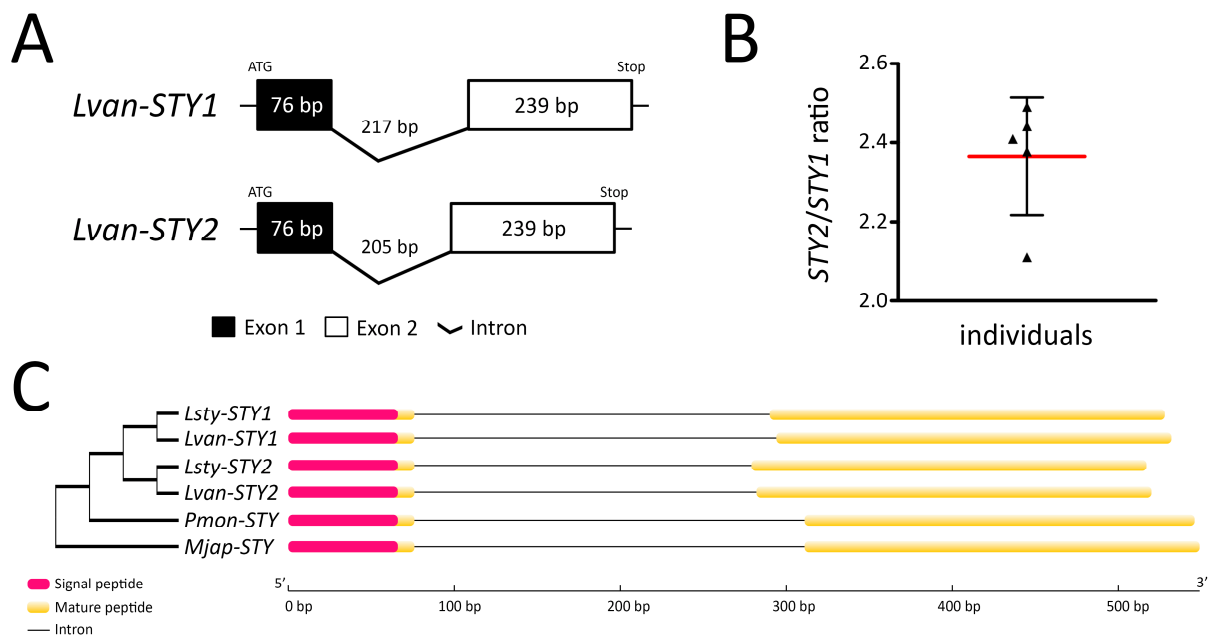
B

Stylicin 1 Stylicin 2

Stylicin 2

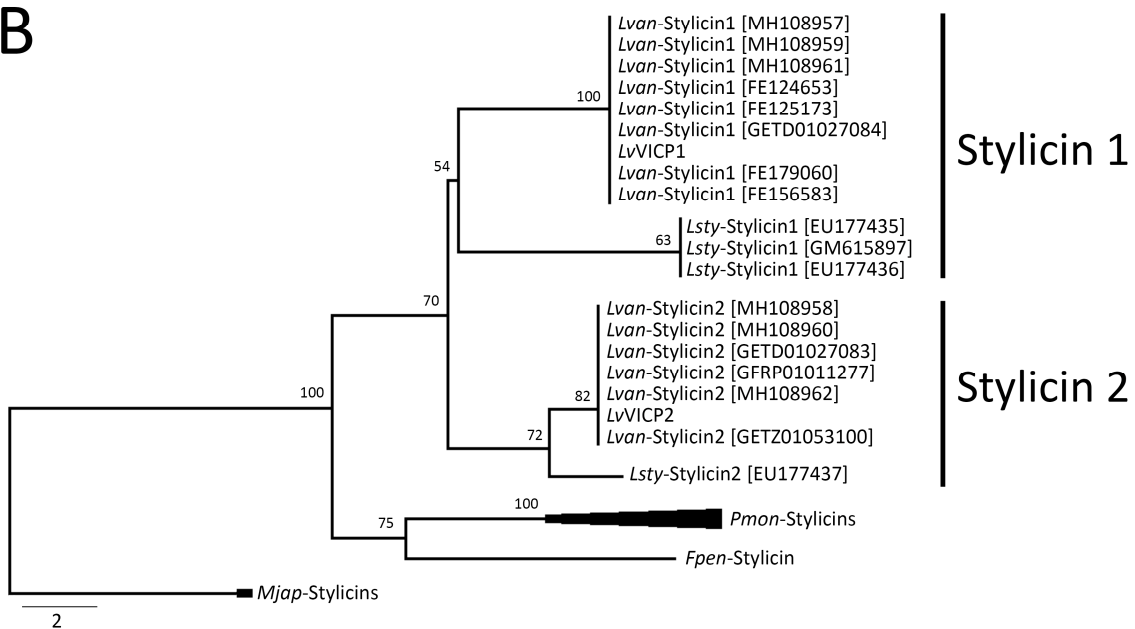
Lvan [MH108958]	MKTYSQSVFVLLVIAIHTQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
Lvan [MH108960]	MKTYSQSVFVLLVIAIHTQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
Lvan [MH108962]	MKTYSQSVFVLLVIAHTSQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
LvVICP2	MKTYSQSVFVLLVIAHTSQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
Lvan [GETD0102708]	MKTYSQSVFVLLVIAIHTQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
Lvan [GFRP010112]	MKTYSQSVFVLLVIAIHTQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
Lvan [GETZ0105310]	MKTYSQSVFVLLVIAHTSQG	SSESPPPGRPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG
Lsty [EU177437]	MKTYSRSVFVLLVIVHTSQG	SSESPPSRLGPLGWEPCCWPCECPCPYDD-ECPKCEEVPAECE-BPDHICDCDPFYHSCLQRGPVCEPCESPIAELIKGGYKG

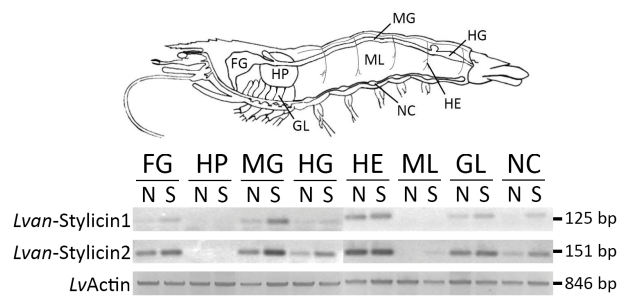
Signal peptide Proline-rich Cysteine-rich

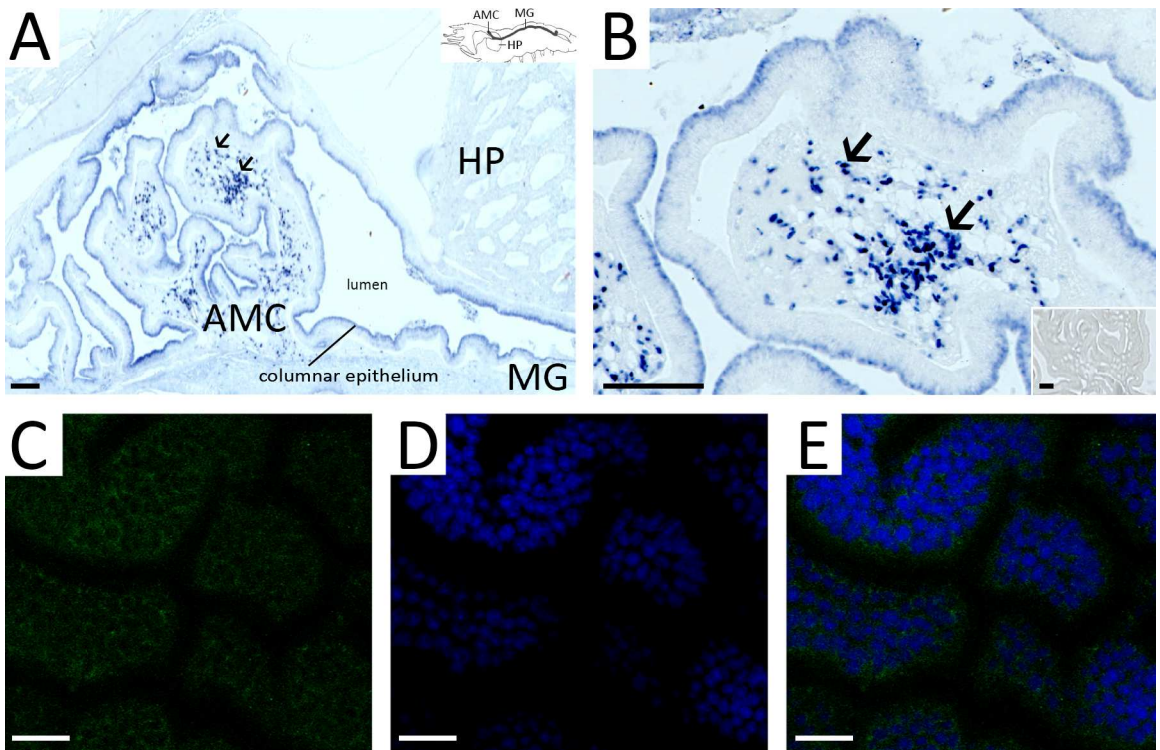


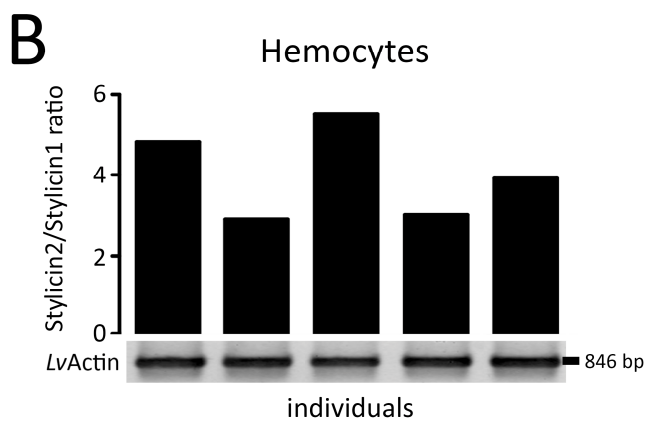
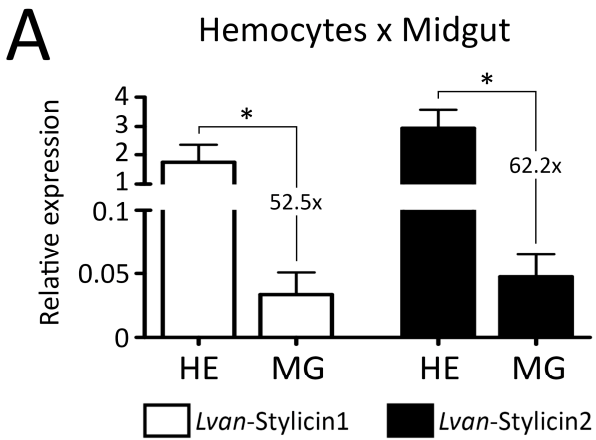
A

<i>Lvan-Stylicin1</i> [MH108957]	SSFSPPRG-PPGNKLPQCPVQECPPCPYDD-ECPRGGFPVQHEVCTDISISCECG-YHSCCOPRPVCPCESPIAELIKKGGYKG
<i>Lvan-Stylicin1</i> [FE179060]	SSFSPPRG-PPGNKLPQCPVQECPPCPYDD-ECPRGGFPVQHEVCTDISISCECG-YHSCCOPRPVCPCESPIAELIKKGGYKG
<i>Lsty-Stylicin1</i> [EU177435]	SSFSPPRG-PPGNGPPCPVQQPCKCPYDDYKCP1CDKFPECEE-CPHISIGCECG-YFSCBOPKPVCBPCESPIAELIKKGGYKG
<i>Lvan-Stylicin2</i> [MH108958]	SSFSPPGR-LPGNEPPCPVQECPPCPYDD-ECPRCEEVPAEE-CPDIHICDCDPFYHSCLQRQPVCBPCESPIAELIKKGGYKG
<i>Lsty-Stylicin2</i> [EU177437]	SSFSPPSR-LPGNEPPCPVQOCPPCPYDD-ECPRGGFPAGEE-CPDIHICDCDPFYHSCLQRQPVCBPCESPIAELIKKGGYKG
<i>Pmon-Stylicin</i> [JZ892895]	SSFYAPSG-PPGERHPCPQLCPVCPG---ECPKCESFPVQHEVCPDIAIAACDOPFYHSCCECRHRVCPPCENPIAELIKKGSYKG
<i>Fpen-Stylicin</i> [GFRTO10057]	SSSFPPWG-PPGWRD1COPQLCPVCPG---ECPKGPFPVQHEVCPDIAIAACDOPFYHSCCECRHRVCPPCENPIAELIKKGSYKG
<i>Mjap-Stylicin</i> [KR063277]	SSFSPAAPLPPGTKHPLSLSCPCP-DE-ECPTCEILPPQEL-OPEIHIICDCDPFHHSQCDQPAACPDPFGSLINKGGYRG

B







Vibrio harveyi