Understanding microRNA Regulation Involved in the Metamorphosis of the Veined Rapa Whelk (Rapana venosa)

Hao Song,*†, Lu Qi,* Tao Zhang,*†, and Hai-yan Wang*†,1

*Chinese Academy of Sciences Key Laboratory of Marine Ecology and Environmental Sciences, Institute of Oceanology, Chinese Academy of Sciences, and 1Laboratory for Marine Ecology and Environmental Science, Qingdao National Laboratory for Marine Science and Technology, 266071, China, 2University of Chinese Academy of Sciences, Beijing 100049, China, and 8College of Fisheries, Ocean University of China, Qingdao 266001, China

ORCID ID: 0000-0003-2197-1562 (H.S.)

ABSTRACT The veined rapa whelk (Rapana venosa) is widely consumed in China. Nevertheless, it preys on oceanic bivalves, thereby reducing this resource worldwide. Its larval metamorphosis comprises a transition from pelagic to benthic form, which involves considerable physiological and structural changes and has vital roles in its natural populations and commercial breeding. Thus, understanding the endogenous microRNAs (miRNAs) that drive metamorphosis is of great interest. This is the first study to use high-throughput sequencing to examine the alterations in miRNA expression that occur during metamorphosis in a marine gastropod. A total of 195 differentially expressed miRNAs were obtained. Sixty-five of these were expressed during the transition from pre-competent to competent larvae. Thirty-three of these were upregulated and the others were downregulated. Another 123 miRNAs were expressed during the transition from competent to postlarvae. Ninety-six of these were upregulated and the remaining 27 were downregulated. The expression of miR-276-y, miR-100-x, miR-183-x, and miR-263-x showed a >100-fold change during development, while the miR-242-x and novel-m0052-3p expression levels changed over 3000-fold. Putative target gene coexpression, gene ontology, and pathway analyses suggest that these miRNAs play important parts in cell proliferation, migration, apoptosis, metabolic regulation, and energy absorption. Twenty miRNAs and their target genes involved in ingestion, digestion, cytoskeleton, cell adhesion, and apoptosis were identified. Nine of them were analyzed with real-time polymerase chain reaction (PCR), which showed an inverse correlation between the miRNAs and their relative expression levels. Our data elucidate the role of miRNAs in R. venosa metamorphic transition and serve as a solid basis for further investigations into regulatory mechanisms of gastropod metamorphosis.

KEYWORDS miRNA metamorphic transition gastropod larval
undertaken by many enterprises since 1992 owing to its economic importance (Yuan 1992). However, large-scale aquaculture of this species has been hampered by difficulties with larval culture during metamorphosis. Moreover, wild veined rapa whelk resources have been declining in China owing to increasing fishing activity. This whelk is considered an invasive species beyond the western Pacific Ocean as a result of unintended world-wide transportation, and it heavily threatens the biomass of local bivalves (Yuan 1992). It was first recorded as an invasive species in the Black Sea during the 1940s (Drapkin 1963). Since then, primarily owing to unintended global transport, R. venosa has become extremely pervasive and has extended its range to Quiberon Bay, France (Mann et al. 2004); Chesapeake Bay (Harding and Mann 1999); Rio de la Plata between Uruguay and Argentina (Pastorino et al. 2000); and The Netherlands’ coastal waters (Nieweg et al. 2005). Its prevalence heavily disrupts native trophic structure and damages endemic bivalve resources. Metamorphosis may control population dynamics; therefore, understanding its mechanism may be useful for aquaculture, resource restoration, and preventing biological invasion of R. venosa.

There are few published reports on R. venosa metamorphosis. A previous study described morphological changes that R. venosa undergoes in the metamorphosis process; the study also indicated that during this process, the diet of R. venosa shifts from phytophagous to carnivorous (Pan et al. 2013). Metamorphosis inducers for this species were also investigated, and it was found that acetylcholine chloride and calcium chloride (CaCl2) were effective and had low toxicity (Yang et al. 2015), suggesting that these compounds may be promising in aquaculture, resource restoration, and preventing biological invasion of R. venosa.

Metamorphosis is the most important developmental event in the molluscan life cycle; however, the characterization and roles of miRNAs in molluscan metamorphosis have not been determined to date. The purpose of the present study was to elucidate the endogenous miRNAs that drive metamorphosis in the veined rapa whelk R. venosa. By sequencing on the Illumina HiSeq 2500 platform, we compared the global expression profiles of small RNAs in precompetent larvae (pre-CL), competent larvae (CL), and postlarvae (PL). In previous studies, we investigated the mRNA global expression profile of whelk metamorphosis (Song et al. 2016d); therefore, in the present study we performed a differentially expressed miRNA-mRNA correlation analysis to elucidate miRNA regulation in whelk metamorphosis. These findings will provide new insights into gastropod metamorphosis and facilitate investigation of miRNA function in a biphasic life cycle in the future.

**MATERIALS AND METHODS**

**Sampling**

Parent R. venosa were collected from their naturally growing areas in Laizhou Bay, China (37°11′4.78″N, 119°41′3.75″E). Parent culture, spawning, larval incubation, and rearing were performed at the Blue Ocean (Laizhou, Shandong, China) according to previously published methods (Pan et al. 2013). Planktonic larvae were cultured in 2.5 × 4 × 1.2 m cement pools at 23.5–25.8°C and a density of 0.3/ml. Ischyris simplex galbana, Chlrella vulgaris, and Platymeris subcordiformis were pooled and provided as a daily diet (2 × 10^6 cells/ml, two times) to the pelagic larvae. Samples from the postlarval stage (three spiral-whorls) stage, the competent larval (four spiral-whorls) stage, and the postlarval (juvenile) stage were collected as three biological replicates, each of which consisted of 40–100 individuals. The samples were inspected under a microscope to ensure that >95% of the individuals were developmentally synchronized. Each replicate was then rinsed with double-distilled H2O and flash-frozen in liquid nitrogen until use.

**Library construction and sequencing**

Total RNA was extracted from an individual intestine using the RNeasyMini Kit (Qiagen, Germantown, MD) according to the manufacturer’s instructions. The quality and concentration of RNA were measured using a NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA). RNA molecules in the size range of 18–30 nt were enriched by polyclayamide gel electrophoresis (PAGE). The 3′ adapters were added and the 36–44 nt RNAs were enriched. The 5′ adapters were then ligated to the RNAs as well. The ligated products were reverse-transcribed by polymerase chain reaction (PCR) amplification. The 140–160-bp PCR products were enriched to generate a cDNA library.
and sequenced using Illumina HiSeq 2500 (Gene Denovo Biotechnology, Guangzhou, China).

**Sequence data analysis**

Raw data were obtained from base calling on the original image. They were cleaned by removing reads containing poly-N or poly-A/T/C/G and 5’ adapter contaminants, and those missing 3’ adapters or insert tags. Low-quality reads were also eliminated. Other RNAs (tags originating from protein-coding genes, repeat sequences, rRNA, tRNA, snRNA, and snoRNA) were deleted after blasting against the RepeatMasker (www.repeatmasker.org/), the GenBank database (http://blast.ncbi.nlm.nih.gov), and the Rfam database (http://sanger.ac.uk/software/Rfam). No miRNA information for rapa whelk is included in miRBase v. 21.0, so the remaining clean reads were aligned to it with ≤2 mismatches to seek all known precursor/mature miRNAs. The miRNAs selected as temporary miRNA references. Clean data were aligned to the highest expression for each mature miRNA family were estimated as temporary miRNA references. Clean data were aligned to them and their expression levels were calculated by summing the read counts aligned with the temporary miRNA database with ≤2 mismatches. The precursors of the identified miRNAs were predicted. Molecules without a hairpin structure were identified as pseudo-miRNAs. The potential novel miRNAs were detected by using MIREAP (http://sourceforge.net/projects/mireap/) with stem-loop structure prediction (Chen et al. 2013).

By pairwise comparison of the miRNA expressions among precompetent, competent and postlarval samples, the differentially expressed miRNAs were identified and estimated. This procedure conforms to the BGI standard as follows: (1) the expression of the miRNA in each sample was normalized to determine the TPM (transcripts per million). The formula TPM = actual miRNA count/total count of clean reads × 10^6 was normalized. (2) The final TPM in each larval stage was calculated by averaging three biological replicates. (3) Fold-change and P-values were calculated based on the normalized expression as described previously (Sun et al. 2017).

**Quantitative real-time PCR**

The total RNA isolated as described above was used in real-time PCR. Briefly, 1 μg of total RNA was reverse-transcribed into cDNA using the One Step PrimeScript miRNA cDNA Synthesis Kit (TaKaRa Bio, Kusatsu, Shiga, Japan), following the manufacturer’s directions. For the miRNA quantitative PCR (qPCR) assay, the primers (Supplemental Material, Table S1) were designed using Primer3 (http://primer3.sourceforge.net/releases.php). The reaction proceeded as follows: 95°C for 3 min; 95°C for 15 sec, 60°C for 25 sec, 72°C for 15 sec for 40 cycles, and a final 20-min extension step at 72°C. Rl28 was chosen as a control gene for internal standardization (Song et al. 2017). The mRNA expression levels were quantified with the SYBR PrimeScript RT-PCR Kit II (TaKaRa Bio) following the manufacturer’s instructions on an Eppendorf Mastercycler ep realplex platform (Eppendorf, Hamburg, Germany). For the miRNA qPCR assay, the reaction proceeded for 60 min at 37°C and 5 sec at 85°C. The CDNA was amplified using real-time PCR and platinum SYBR Green qPCR SuperMix-UDG (Invitrogen, Carlsbad, CA) with miRNA-specific forward and reverse primers (Table S1). The 5.8 S rRNA was used as an internal reference gene to normalize the data (Zheng et al. 2016). The amplification products were detected by melting curve and gel electrophoresis to ensure primer efficiency and PCR specificity. The relative expression levels of miRNA and mRNA were estimated by the 2−ΔΔCT method. All data were presented as means ± SE (N = 3). Statistical significance was analyzed using SPSS v. 18, with P < 0.05 being considered significant.
Target gene prediction and correlation analysis of miRNA-mRNA

The 3’ UTRs from the rapa whelk transcriptome assembly (Song et al. 2016c,d) were used as a reference database to predict the target genes with RNA hybrid v. 2.1.2, svm_light v. 6.01, Miranda v. 3.3a, and TargetScan v. 7.0. The targeting criteria were: (1) no mismatch is allowed between 2 and 8 nt on the 3’ end; (2) the G–U matching number cannot be >3; and (3) the minimum free energy (MFE) of the miRNA/target duplex should be >75% of the MFE of the miRNA bound to its perfect complement. The target genes of differentially expressed miRNAs were mapped to the gene ontology (GO) database (http://www.geneontology.org/) and the KEGG (Kyoto Encyclopedia of Genes and Genomes; http://www.genome.jp/kegg/) pathways for GO and KEGG analyses. GO and pathway terms with \( P < 0.05 \) (Bonferroni’s correction) were considered statistically significant. The differentially expressed mRNA identified in a previous study (Song et al. 2016d) and the differentially expressed miRNAs were integrated to analyze the key miRNA-target pairs. Only the inversely correlated miRNA-target pairs with MFE \( \leq -18 \) were screened.

Data availability

Raw sequencing data were submitted to the GEO (Gene Expression Omnibus) database with accession No. GSE102631. Supplemental materials include the details of primers for qPCR assays (Table S1), the differentially expressed miRNAs during metamorphosis development (Table S2), and miRNA-target pairs of differentially expressed miRNAs (Table S3).

RESULTS

MicroRNA library construction

To identify the miRNAs differentially expressed during metamorphosis in rapa whelk, nine small RNA libraries (precompetent larvae: Pre-CL 1, Pre-CL 2, and Pre-CL 3; competent larvae: CL 1, CL 2, and CL 3; postlarvae: PL 1, PL 2, and PL 3) were constructed using Illumina sequencing. A dataset consisting of \( \approx 85,000,000 \) reads (ranging from 8,235,539 to 10,159,366 among the samples) was obtained after trimming the adapter sequences (Table 1). A BLAST run against the NCBI GenBank, the RepeatMasker, and the Rfam database identified 19,494 (8.75%) to 18,982 (11.77%) unique small RNAs as rRNA, 229 (0.11%) to 635 (0.19%) as snRNA, 28 (0.01%) to 68 (0.03%) as snoRNA, and 1284 (0.62%) to 3619 (1.12%) as tRNA (Table 1). After removing these small RNAs, the remaining RNAs were further analyzed to identify rapa whelk miRNAs against miRBase v. 21.0. A total of 19,878 (12.33%), 20,976 (11.31%), and 18,263 (9.69%) unique known miRNAs were sought in precompetent larvae, 21,033 (9.44%), 17,913 (9.51%), and 16,210 (7.85%) in competent larvae, and 20,573 (6.36%), 24,996 (7.57%), and 22,325 (6.56%) in postlarvae. There were 212 (0.13%), 211 (0.11%), and 209 (0.11%) unique novel miRNAs found in precompetent larvae, 192 (0.09%), 191 (0.10%), and 181 (0.09%) in competent larvae, and 173 (0.05%), 205 (0.06%), and 207 (0.06%) in postlarvae.

Different expression profiles of miRNAs

The differentially expressed miRNAs were identified by \( t \)-test with a fold change >2 and \( P < 0.05 \). A total of 195 miRNAs were obtained, including 33 upregulated and 32 downregulated during the transition from precompetent to competent larvae, and 96 upregulated and 27 downregulated during metamorphosis (Figure 1 and Table S2). Table 2 lists 39 differentially expressed miRNAs with the following criteria: average TPM \( >10 \) (in nine samples), \( \log_{2} \text{Ratio} > 2 \) or \( < -2 \) and \( P < 0.01 \) for \( \geq 1 \) comparison among the groups. Eleven miRNAs showed a >20-fold difference in gene expression for \( \geq 1 \) comparison among the groups. These molecules included miR-242-x, novel-m0052-3p, miR-276-y, miR-100-x, miR-183-x, miR-263-x, miR-99-x, miR-37-y, miR-36-y, miR-1175-y, and miR-125-x. These may have important roles in regulating metamorphosis-associated gene expression. Specifically, miR-276-y, miR-100-x, miR-183-x, and miR-263-x showed a >100-fold change, while miR-242-x and novel-m0052-3p expression levels showed a >3000-fold change.

GO and KEGG pathway enrichment of miRNA target genes

The putative target genes of differentially expressed miRNAs were identified using TargetScan software based on rapa whelk transcriptome libraries (Song et al. 2016c). GO analysis was then used to predict enriched functional groups (\( P < 0.05 \) (Figure 2). “Cellular process (10,917 genes),” “metabolic process (9669 genes),” and “single-organism process (8590 genes)” were the top three enriched items in the biological process, whereas “binding (10,297 genes),” “catalytic activity (8104 genes),” and "biological process (1284 (0.62%) to 3619 (1.12%) as tRNA (Table 1). After removing these small RNAs, the remaining RNAs were further analyzed to identify rapa whelk miRNAs against miRBase v. 21.0. A total of 19,878 (12.33%), 20,976 (11.31%), and 18,263 (9.69%) unique known miRNAs were sought in precompetent larvae, 21,033 (9.44%), 17,913 (9.51%), and 16,210 (7.85%) in competent larvae, and 20,573 (6.36%), 24,996 (7.57%), and 22,325 (6.56%) in postlarvae. There were 212 (0.13%), 211 (0.11%), and 209 (0.11%) unique novel miRNAs found in precompetent larvae, 192 (0.09%), 191 (0.10%), and 181 (0.09%) in competent larvae, and 173 (0.05%), 205 (0.06%), and 207 (0.06%) in postlarvae.

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“transporter activity (1732 genes)” participated in molecular function. “Cell (6406 genes),” “cell part (6406 genes),” “membrane (5487 genes),” “organelle (4219 genes),” and “macromolecular complex (4059 genes)” were highly represented in the cellular component categories.

Enriched metabolic and signal transduction pathways were identified and are listed in Figure 3. Eight significantly enriched pathways for target genes (Q < 0.05) involved in “TNF signalling pathway,” “SNARE interactions in vesicular transport,” “Glycosylphosphatidylinositol (GPI)-anchor biosynthesis,” “Nicotinate and nicotinamide metabolism,” “Ubiquitin-mediated proteolysis,” “Phosphonate and phosphinate metabolism,” “Pyrimidine metabolism,” and “Sulphur relay system” were screened (Figure 3).

Selection of miRNA-target pairs and qPCR validation

The aforementioned mRNA expression profiling data from the same metamorphosis sampling stages (Song et al. 2016c) were used to perform association analyses along with the current miRNA profiling. The differentially expressed miRNAs detected in the present study were used to select the miRNA-target pairs expressed in metamorphosis (Table S3). The miRNAs negatively regulate the expression of their target mRNAs either by translation inhibition or by mRNA degradation. In previous studies (Song et al. 2016b,c), the miRNAs/proteins involved in “ingestion and digestion,” “cytoskeleton and cell adhesion,” and “apoptosis” were thought to have important roles in driving metamorphosis. We identified 20 key miRNA-target pairs potentially implicated in these aspects of whelk metamorphosis (Table 3). For example, in “ingestion and digestion,” we found that let-7-y potentially regulates the SARP-19 precursor, conotoxin C14.12, and the exogulcanase XynX genes. Therefore, a single miRNA may regulate multiple target genes during metamorphosis. The miR-1175-x targets the cysteine-rich secretory protein gene, and miR-2001-x targets endo-1,4-β-xylanase. The gene mir-71-x regulates the β-1,4-xylanase and membrane metalloendopeptidase-like 1 genes. In “cytoskeleton and cell adhesion,” tektin-3, dynein heavy chain 8 (axonemal), and dynein intermediate chain 2 (ciliary), all of which are the main structures of velum cilia, were regulated by miR-5106-y, miR-87-y, and miR-315-x, respectively. Novel-m0020-5p determines the expression of
apoptosis 2 inhibitor genes, and miR-276-y regulates the caspase-3 gene. These two mRNAs are both important for programmed cell death during metamorphosis.

Real-time PCR analysis was performed on nine key targeted miRNA-mRNA pairs (Figure 4) to validate and identify the metamorphosis-related miRNAs in rapa whelk. The results showed that the miRNAs...
were consistent with the overall trend in high-throughput sequencing. For each of the nine pairs, there was an inverse correlation between the expression levels of miRNA and miRNA. For example, the let-7-y miRNA decreased during the transition from precompetent to postlarval, whereas its target mRNAs (c112229_g1 SARP-19 precursor, c124801_g1 Conotoxin Cl14.12, and c150903_g1 Exoglucanase XynX) significantly increased.

**DISCUSSION**

In this study, we investigated the miRNA expression profile during the metamorphosis of the rapa whelk using high-throughput sequencing. In total, nine libraries were constructed and 85,000,000 reads were obtained. These results will augment information on the small RNA genome of rapa whelk and provide a basis for miRNA regulation during metamorphosis. A total of 195 miRNAs was significantly differentially expressed among three larval stages (precompetent, competent, and postlarval stages) and targeted thousands of genes. This result was expected since metamorphosis is the most complicated developmental event of the life cycle and entails considerable structural, physiological, and behavioral changes.

To obtain an insight into the possible functions of the differentially expressed miRNAs involved in metamorphosis, we performed GO and KEGG pathway enrichment analyses on their predicted target genes. The significantly enriched GO terms “biological adhesion,” “cell aggregation,” “cellular component organisation or biogenesis,” “localisation,” “developmental process,” “signalling,” “immune system,” and “response to stimulus” are primarily associated with development, gene expression, immunity, and the cell cycle. Metamorphosis is a complex process and includes tissue remodeling, cell migration, differentiation, proliferation, and others (Jackson et al. 2005). A total of eight significantly enriched pathways were observed. The “TNF signalling” pathways were enriched because they trigger apoptosis, and old organs such as the velum are degenerated by programmed cell death. “SNARE interactions in vesicular transport” were enriched because the nervous system mediates metamorphosis in many marine invertebrates (Voronezhskaya 2004; Gifondorwa and Leise 2006) and SNARE participates in vesicle docking, priming, fusion, and the synchronization of neurotransmitter release into the synaptic cleft during neurosecretion (Shi et al. 2011). SNAREs also play a crucial part in the autophagy required for velum degradation and reabsorption during molluscan metamorphosis. “Nicotinate and nicotinamide metabolism,” “Ubiquitin mediated proteolysis,” “Phosphonate and phosphinate metabolism,” “Pyrimidine metabolism,” and “Sulphur relay system” were also affected because of the tissue remodeling and energy redistribution that occur during metamorphosis.

Table 2 lists 39 differentially expressed miRNAs with striking changes. The miR-242-x expression level steadily decreases as precompetent larvae develop into postlarvae. Its expression level decreased by >3000 fold after metamorphosis. Therefore, the miR-242 family may have important roles.
rubio and belles 2013). in the fruit miRNAs in this species are of great interest. In signi three miRNAs were found at low levels in competent larvae but rose to a high level in postlarvae. The function of this miRNA remains as yet un-

than 0.01 in the competent larvae and rose sharply to 3974.4 after metamorphosis. There was a similar trend for miR-125-x. Both miR-100-x and miR-125 are believed to participate in cell migration. Low levels of miR-100 and miR-125 may promote hepatocellular carcinoma metastasis (Rubio and belles 2013). Rapa whelk metamorphosis is accompanied by high levels of cell death, proliferation, and tissue remodeling, thus implicating that let-7 participates in digestive system changes during whelk metamorphosis, whereas those of its target miRNAs like SARP-19, conotoxin, and exogluccanase continuously declined. The SARP-19 gene was expressed highly in the gastropod larval digestive gland and can be sensitive to metamorphic cues (he et al. 2014). Conotoxin is a group of neurotoxic peptides isolated from Conus venom. High conotoxin levels in R. venosa pelagic larvae indicated that this life stage in R. venosa is homologous with that of Conus. In the rapa whelk, however, it degenerated after metamorphosis (song et al. 2016b,c). Exogluccanase, an important digestive enzyme in whelk pelagic larvae, sharply decreased when the whelk become carnivorous after metamorphosis. The digestion-related genes were negatively regulated by let-7-y, which implies that let-7 participates in digestive system changes during whelk metamorphosis. The let-7 miRNA also participates in metamorphosis in rapa whelk metamorphosis. In Caenorhabditis elegans, mir-242 and mir-793 target the Argonaute protein ALG-1, which controls the RNA interference process involved in developmental timing (grishok et al. 2001). The expression level of novel-m0052-3p in precompetent larvae remained at an average TPM of 8.17. On the other hand, the average TPM decreased to 0.01 in the competent larvae and rose sharply to 36.61 in the postlarvae. The function of this miRNA remains as yet un-known and, to our knowledge, no relevant studies on it have been published to date. The TPM level of mir-100-x in the precompetent larvae was 59.29. It decreased to 23.12 in the competent larvae but sharply increased to 4006.4 after metamorphosis. As stated in previous studies (Song et al. 2016b,c), the miRNAs/proteins involved in ingestion and digestion, cytoskeleton and cell adhesion, and apoptosis may also have important roles in driving metamorphosis. Twenty key miRNA-target pairs implicated in these processes were identified, and nine of them were further studied by real-time PCR. The let-7 miRNA expression level was low in competent larvae. This developmental stage involves considerable amounts of programmed cell death because the velum must degenerate. Therefore, the levels of miR-9 must be high after metamorphosis in order to suppress further apoptosis.


table 3 coexpression of 20 key miRNA-targets

| No. | miRNA ID | Method | Average miRNA TPM | Target Gene ID | Average mRNA FPKM |
|-----|----------|--------|-------------------|---------------|------------------|
| 1   | let-7-y  |        | 8.32              | c112229_g1    | 252.36           |
| 2   | let-7-y  |        | 8.32              | c124801_g1    | 9144.35          |
| 3   | let-7-y  |        | 8.32              | c150903_g1    | 69.11            |
| 4   | mir-1175-x |      | 76.60             | c119967_g1    | 19.39            |
| 5   | mir-2001-x |      | 87.40             | c154241_g2    | 10.27            |
| 6   | mir-87-y  |        | 1480.69           | c124801_g1    | 9144.35          |
| 7   | mir-981-y |        | 3915.01           | c137870_g1    | 1261.55          |
| 8   | mirror-71-x |      | 12183.40          | c156902_g1    | 29.13            |
| 9   | mirror-71-x |      | 12183.40          | c156902_g1    | 29.13            |
| 10  | mirror-71-x |      | 12183.40          | c156902_g1    | 29.13            |
| 11  | mir-S106-y |        | 0.68              | c155866_g1    | 833.87           |
| 12  | mir-87-y  |        | 1480.69           | c157287_g2    | 69.54            |
| 13  | mir-315-y |        | 25652.61          | c154991_g1    | 87.45            |
| 14  | mir-283-x |        | 489.56            | c137644_g1    | 1.61             |
| 15  | mir-263-x |        | 57623.55          | c146951_g1    | 46.76            |
| 16  | novel-m0020-5p |      | 33.74             | c151900_g1    | 4.23             |
| 17  | mir-276-y |        | 44.03             | c135194_g1    | 0.78             |
| 18  | mir-92-x  |        | 67.63             | c135194_g1    | 0.78             |
| 19  | mir-183-x |        | 51432.32          | c105989_g1    | 76.77            |
| 20  | mir-216-x |        | 43564.02          | c105989_g1    | 128.37           |

Pairs with asterisks were further analyzed by real-time PCR.
in many other animals. In Drosophila, for example, the loss of let-7 and miR-125 may delay the terminal cell-cycle exit in the wing and the maturation of neuromuscular junctions (NMJs) in the adult abdominal muscles. The maturation rate of abdominal NMJs was governed by let-7 during metamorphosis by regulating the expression of the ab gene (Caygill and Johnston 2008). In the silkworm Bombyx mori, let-7 regulates molting and metamorphosis. Decreased let-7 expression in the silkworm could increase the expression of its target genes FTZ-F1 and Eip74EF (key regulatory factors in the ecdysone pathway) and cause developmental arrest during the larval–larval and larval–pupal transitions (Ling et al. 2014). The development-related miR-276 may inhibit apoptosis in shrimp hemocytes (Yang et al. 2012). In this study, miR-276 was found to be significantly downregulated in competent larvae, whereas the caspase-3 gene was upregulated. Therefore, miR-276 may regulate apoptosis by targeting the caspase-3 gene. Coexpression studies of key miRNA targets revealed their potential roles in whelk metamorphosis, but the mechanisms by which these miRNAs regulate this developmental process have not been fully explored yet.

In conclusion, the present study provides the first global view of the changes in miRNA that occur during rapa whelk metamorphosis. A total of 195 miRNAs were significantly differentially expressed and their target mRNAs were identified. These molecules are responsible for morphological and functional changes in organs. Some miRNAs involved in ingestion and digestion, cytoskeleton and cell adhesion, and apoptosis during metamorphosis are of great interest and were listed and validated by real-time PCR. These results will provide a basis for understanding the molecular mechanisms involved in the regulation of gastropod metamorphosis.

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