

RESEARCH ARTICLE

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Deep sequencing of small RNA libraries reveals dynamic regulation of conserved and novel microRNAs and microRNA-stars during silkworm development

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Abstract

Background: In eukaryotes, microRNAs (miRNAs) have emerged as critical regulators of gene expression. The Silkworm (*Bombyx mori* L.) is one of the most suitable lepidopteran insects for studying the molecular aspects of metamorphosis because of its large size, availability of mutants and genome sequence. Besides, this insect also has been amply studied from a physiological and biochemical perspective. Deep sequencing of small RNAs isolated from different stages of silkworm is a powerful tool not only for measuring the changes in miRNA profile but also for discovering novel miRNAs.

Results: We generated small RNA libraries from feeding larvae, spinning larvae, pupae and adults of *B. mori* and obtained ~2.5 million reads of 18-30 nt. Sequence analysis identified 14 novel and 101 conserved miRNAs. Most novel miRNAs are preferentially expressed in pupae, whereas more than 95% of the conserved miRNAs are dynamically regulated during different developmental stages. Remarkably, the miRNA-star (miR*) of four miRNAs are expressed at much higher levels than their corresponding miRNAs, and their expression profiles are distinct from their corresponding miRNA profiles during different developmental stages. Additionally, we detected two antisense miRNA loci (miR-263-S and miR-263-AS; miR-306-S and miR-306-AS) that are expressed in sense and antisense directions. Interestingly, miR-263 and miR-306 are preferentially and abundantly expressed in pupae and adults, respectively.

Conclusions: We identified 101 homologs of conserved miRNAs, 14 species-specific and two antisense miRNAs in the silkworm. Our results provided deeper insights into changes in conserved and novel miRNA and miRNA* accumulation during development.

Background

Transcriptional regulation alone is insufficient to ensure tight control of gene expression in specific cells or tissues. Recently discovered microRNA (miRNA)-directed post-transcriptional regulation can provide an efficient fine-tuning of target gene expression in certain cell or tissue types and, thus, coordinate the spatial and temporal control [1,2]. Furthermore, miRNA-guided suppression of the target genes can be quickly relieved, and

such reactivation is faster than transcriptional activation of a genomic locus; thus, miRNAs can act as reversible regulators [1]. Because of the versatility, miRNAs have evolved as a major class of gene-regulatory molecules critical for diverse biological processes such as cell proliferation, differentiation, apoptosis, stress response, tumorigenesis, diabetes and heart failure in eukaryotes [3-11].

In animals, most miRNA genes are transcribed by RNA polymerase II, yielding transcripts called primary miRNAs (pri-miRNAs), which are initially processed by a complex containing Drosha and then by Dicer-1 to excise miRNA:miRNA-star (miR:miR*) duplexes. One

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strand of the duplex (miR) is more stable and preferentially incorporated into an RNA-induced silencing complex (RISC). The miRNA then guides the RISC to regions of complementarity in the target site, where it downregulates the gene expression, often by blocking protein production or by degrading the target mRNA [12-16].

Insect metamorphosis is a complex, highly conserved, and strictly regulated process of developmental events. During metamorphosis, diverse morphological, physiological, biochemical and molecular events result in distinct changes such as cell proliferation, programmed cell death, tissue remodeling and cell migration [17]. The silkworm is an ideal model for studying metamorphosis in holometabolous insects, because of its large size, the availability of mutants with nearly fully sequenced genome. Additionally, this insect has been amply studied from a physiological and biochemical perspective [18]. Many agricultural pests belonging to Lepidoptera cause economic damage to commercial crops. Thus, molecular studies focusing on silkworm metamorphosis should provide better understanding of insect gene regulation and novel targets for pest control.

Thus far, miRNAs cataloging in insects is primarily carried out in *Drosophilid* species, and many miRNAs were discovered both by direct cloning [19-23] and by bioinformatic methods [24,25]. Besides Drosophila, mosquito (A.gambiae) [26,27], honey bees (A.mellifera) [28,29], red flour beetle [30] and locusts [31] are some of the insect species, in which miRNAs has been identified. Most existing studies of small RNAs in the silkworm have focused on identification of miRNAs using computational strategies [32-35]. Such studies can identify conserved miRNAs but not species-specific ones. Recently cloning of miRNAs in the silkworm has been reported [36], in which Zhang and co-authors analyzed 95,184 unique small RNA reads and annotated 354 of them as miRNAs; none were based on miR* sequences. Surprisingly, 253 of these reported miRNAs (>70%) were represented by single reads, and an additional 51 miRNAs were represented by two reads in the library. One of the most important criteria for annotating novel miRNAs is cloning their miR* sequences and this becomes even more important for annotating speciesspecific miRNAs. We have recovered most of the sequences (>90%) reported by Zhang et al., [23] multiple times in multiple libraries, yet could not annotate them as miRNAs due to the lack of miR* support. Furthermore, Zhang et al. [36] did not measure changes in miRNA abundance at different developmental stages, because RNA from various stages was pooled for library construction in their study.

Deep sequencing of small RNAs can be used to reliably measure modest changes in miRNA abundance

among different samples; such changes are unlikely to be identified by sequencing low numbers of clones (i.e., traditional small RNA library sequencing) or hybridization-based methods such as small RNA blot and miRNA array analyses. The deep sequencing study is also well suited for the discovery of species-specific miRNAs expressed at low abundance. Indeed, in this study, by using deep sequencing we uncovered 15 novel miRNAs that appear to be silkworm-specific. Of the 101 conserved miRNAs identified in this study, most are dynamically regulated during different developmental stages of B. mori. Interestingly, the abundance of miR* species of several conserved miRNAs was greater than that of their miRNAs, and their levels varied greatly at various stages. We also discovered two antisense miR-NAs in the silkworm. This study provides deeper insights into dynamic regulation of the conserved and novel miRNA and miRNA* levels during different developmental stages and suggests that both universal and silkworm-specific miRNAs play important roles in expression regulation.

Results

Insect metamorphosis is a highly complex and integrated set of developmental processes coordinated by the action of hundreds of genes [17]. Because miRNAs are important gene regulators, we monitored the changes in miRNA expression during larval-pupal, and adult stages of the silkworm. We constructed two small RNA libraries, one from feeding larvae (4th and 5th instars) and the other from spinning larvae (Figure 1a). Insects from these two stages differ significantly in physiology. 4th and 5th instar larvae are voracious eaters, whereas spinning larvae do not consume food, instead, they oozes silk and spin cocoon. Besides, the usage of spinning larvae may help to identify transiently regulated miRNAs (consequently, miRNA-mediated processes) during the larval-to-pupal molt. We constructed two additional libraries, from pupae and moths (Figure 1a). Pupae undergo complex histolysis and histogenesis, and their physiology and metabolism are different from that of larvae and adults. The adult library may reveal information on miRNA-mediated regulation of reproduction. While constructing the libraries, we used a barcoded 5' adapter for each of the four RNA samples. This helped us sequence these libraries together to a reasonable depth using the sequencing-by-synthesis (SBS) sequencing technology (Illumina inc.), yet the sequences can be traced back to their original stages.

A dataset of about 2.5 million reads (1 147 753, 599 279, 289 303 and 463 800 reads from the feeding larvae, spinning larvae, pupae, and moths, respectively) ranging from 18 to 30 nt was obtained after trimming the adapter sequences. A significant number of reads (~300,000)

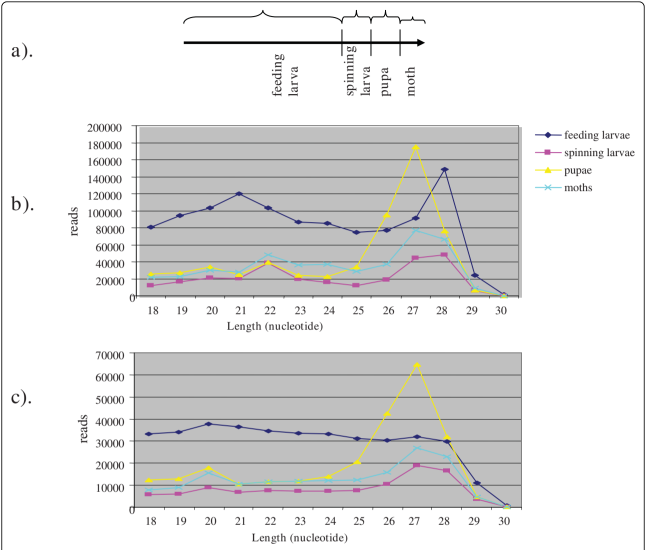


Figure 1 Length distribution and abundance of small RNAs in silkworm. a) Schematic representation of the silkworm life cycle beginning with the feeding larval stage. Abundance of each size class of small RNAs based on nucleotide (nt) length in feeding larvae, spinning larvae, pupae and moths plotted using total small RNA reads (b) and unique reads (c).

lacking the bar-code sequences were eliminated from the analysis because these could not be assigned to the source libraries. From the size distribution of total reads, we found a distinct bimodal distribution, with a peak around 20-22 nt representing miRNAs and another distinct peak around 26-29 nt mostly representing longer piRNA-like small RNAs in all four stages (Figure 1b and 1c). Size-dependent bimodal distribution of small RNAs was also evident, with unique small RNAs plotted from spinning larvae, pupae and moths but not from feeding larvae (Figure 1b and 1c). A peak for the 26 to 29-nt size class with total reads but not unique reads specifically in feeding larvae suggests that these small RNAs are more diverse in larvae and thus not enriched for unique small RNAs. Overall, the peak

represented by the 26- to 29-nt size class was greater than that for the 20- to 22-nt size class and suggested that the longer-sized small RNAs are more abundant in all four stages examined. Notably, the peak represented by the long (26-29 nt) unique small RNAs was much greater in the pupal stage (Figure 1c). The greater abundance of unique small RNAs of long size class in pupae implies a critical role for these small RNAs in the pupal stage.

Of 2.5 million total reads, 1,581,810 could be mapped to the available silkworm genome. We used the latest version of silkDB v2.0 to map and analyze these small RNAs. After eliminating redundant sequences from the dataset, only unique sequences represented by 739,119 reads were analyzed further (Table 1). Subsequent

	Feeding larvae	Spinning larvae	Pupae	Moth	Total
non-codingRNAs (rRNA, tRNAs etc.,)	558345	167369	94118	169508	989340
conserved and novel miRNAs	43278	70143	67256	69913	250590
messenger RNAs	101508	27736	9577	13564	152385
mitochondrial RNAs	7356	951	231	201	8739
repetitive elements	147327	40034	16522	30965	234848
subtotal	832966	293424	182808	272612	1581810
cannot be mapped to the available genome	314787	305855	106495	191188	918325
Total	1147753	599279	289303	463800	2500135
Unique small RNAs	350095	245339	101669	150197	739119

sequence analysis eliminated reads corresponding to rRNAs, tRNAs, snRNAs and snoRNAs and repetitive elements, assuming that these are degradation products [37-39]. The unique dataset with read counts was used to identify conserved and novel miRNAs in silkworm.

Novel miRNAs in silkworm

The appearance of both the miRNA and its corresponding miR* in a dataset would provide compelling evidence for annotation of a novel miRNA. Accordingly, we have annotated 14 small RNAs as novel miRNAs in the silkworm by identifying both miR and miR* sequences in our libraries (Table 2). Using their precursor sequences, we were able to predict fold-back structures for these novel miRNAs (Figure 2). miR* sequences were not represented in our dataset for two small RNAs (bmo-mir-2733e-1 and bmo-mir-2733f) but were annotated as novel miRNAs because these belong to a novel miRNA family with seven members (Table 2). Additionally, we recovered several dozens of unique small RNAs that resemble miRNAs on the basis of sequence characteristics (begin with 'U', 20-23 nt long, and with a predicted fold-back structure for their precursors). Because their miR* sequences were not detected, these were not annotated as miRNAs for now. Retrieval of relatively high number of reads from multiple libraries and appearance of miR*, coupled with predicted fold-back structure for the precursor, strongly supported that these mature miRNAs are processed from hairpin structures in the silkworm. Since, we were unable to find homologs for these miRNAs in related insects or other animal species; we annotated them as species-specific miRNAs.

Sequence alignment indicated that a new miRNA family represented by seven members (bmo-mir-2733a-2, bmo-mir-2733b, bmo-mir-2733d, bmo-mir-2733e-1, bmo-mir-2733f, bmo-mir-2733 g, and bmo-mir-2733 h) was identified in this study (Figure 3a). The seed region (2-8 nt), which is critical for target recognition, is identical in all these seven members, and the sequence conservation extends beyond the seed region. Furthermore, the miR*, for five of them, is also highly

conserved (Figure 3b). The miRNAs, bmo-mir-2733a-2 and bmo-mir-2733 g do not display sequence conservation except in the mature miRNA region. These two miRNAs also display differences with respect to miRNA organization within the hairpin structures; the miRNAs are located in the 5' arm of the hairpin structure, whereas the other five are located in the 3' arm of the predicted fold-back structure (Figure 2). Within an miRNA family, the location (organization) of mature miRNA in the fold-back structure is well conserved [40]. Thus, miRNA organization for these two novel miRNAs appears to be atypical.

Changes in expression profile of the species-specific miRNAs were analyzed based on the normalized read abundance in the four different developmental stages. Two novel miRNAs (bmo-mir-2763 and bmo-mir-3001) were present at low levels in larvae relative to other stages (Table 2). The miRNA, bmo-mir-2766 was ubiquitously expressed in all four developmental stages with its highest level found in pupae. Similarly, eight novel miRNAs (sRNA086545, bmo-mir-2733a-2, bmomir-2733 g, bmo-mir-2733e-1, bmo-mir-2733f, bmomir-2733b, bmo-mir-2733d and bmo-mir-2733 h) were highly expressed in pupae relative to the other stages. Three novel miRNAs (bmo-miR-2998, bmo-mir-2999 and bmo-mir-2763) were most abundantly expressed in the moth stage. Individual counts of the miRNA reads varied considerably, indicating a differential regulation of expression for the novel miRNA family members (Table 2). MicroRNA, bmo-mir-2733a-2 was the most abundantly expressed, followed by bmo-mir-2733 g (Table 2). Taken together, most novel miRNAs were expressed in a stage-specific manner in silkworm. Consistent with these results, genes essential for insect development and metamorphosis are predicted targets for novel miRNAs and miRNA* in silkworm (Additional file 1). We have predicted over 1100 genes as targets for the newly identified silkworm-specific miRNAs and their star species (Additional file 1). These findings suggest an important regulatory role for novel miRNAs in silkworm development.

Table 2 Identification of novel silkworm-specific miRNAs based on sequencing miRNAs and miRNA_Stars in different
developmental stages.

miRNA	Sequence of mature miRNA	Feeding larvae		Spinning larvae		Pupae		Moths		Remarks
		miR	miR*	miR	miR*	miR	miR*	miR	miR*	_
bmo-mir-2998	AAGAACAGGAUGAGGUAGAUAAA	26	1	9	0	10	0	88	1	
bmo-mir-375	ACCCGAGCGGUCUGAGCAAACU	30	6	4	2	25	11	11	3	
bmo-mir-2766	UCAGUCUUGUCGAAUGGUGGGU	2205	284	2657	166	4588	341	2328	133	
bmo-mir-2999	CUGCGACGGACUAGACGCGCA	20	2	3	0	16	0	107	0	
bmo-mir-2763	AUAUUAUGCUCAUUUCUUUGGAU	15	0	49	0	39	1	199	0	
bmo-mir-2733e-1	UCACUGGGAAUGUAAUAGCUAU	1	0	4	0	273	0	1	0	Family
bmo-mir-2733f	UCACUGGGUAUGUAAUGACAGU	1	0	0	0	83	0	1	0	Family
bmo-mir-2733 g	UCACUGGGUGCAUGAAGAUUG	2	0	0	0	629	2	1	0	Family
bmo-mir-2733a-2	UCACUGGGUGCAUGAUGAUUG	16	0	6	0	2718	2	93	0	Family
bmo-mir-2733b	UCACUGGGUGCGUGAUGAUUGU	0	0	0	0	134	3	0	0	Family
bmo-mir-2733d	UCACUGGGUGUAUGAAGAUUG	0	0	2	0	392	1	5	0	Family
bmo-mir-2733 h	UCACUGGGUGUAUGAUGAUUG	0	0	1	0	163	2	1	0	Family
bmo-mir-3000	CUGCGCUUAGAUGAAGACACUA	53	1	7	0	6	0	6	0	
bmo-mir-3001	UAAGUUGAAAGAAUUGUAGAUUUUGA	4	7	7	8	12	2	14	14	

Overall miRNA abundance is dynamically regulated during successive developmental stages of silkworm

The proportion of miRNAs to the overall small RNA population varied greatly in different developmental periods. miRNAs accounted for 3.7%, 11.70%, 23.24%, and 15.07% of the total small RNA component from feeding larvae, spinning larvae, pupae and moths, respectively, indicating that overall miRNA abundance changes greatly between any two successive developmental stages of *B. mori.*

The abundance of a given miRNA relative to the overall miRNA abundance in each library was calculated as number of reads of a given miRNA/total number of reads of all miRNAs. Previous reports indicated that in diverse animal species, evolutionarily conserved miRNAs are often highly expressed [38,41,42]. The top three abundant miRNAs of the silkworm are miR-1 (30%), miR-8 (11%) and miR-306 (11%) in the feeding larval stage; miR-1 (40%), miR-8 (18%), and miR-276a (15%) in the spinning larval stage; miR-276a (27%), miR-1 (16%) and miR278 (8%) in the pupal stage; miR-263a (18%), miR276a (16%) and miR-1 (12%) in the adult stage (Additional file 2). Thus, depending on developmental stage, the most abundant miRNA (one individual miRNA) occupies between 18% and 40% of the overall miRNA populations. Furthermore, the top three abundant miRNAs in any given stage cover between 46% and 73% of the overall miRNA populations. This indicates that the most abundantly expressed miRNAs in each stage are different and the levels vary greatly between stages. Interestingly, miR-1 and miR-276a are two of the three most abundant miRNAs in all four stages.

Temporal regulation of conserved miRNAs during silkworm development

In this study, we found evidence for the expression of 101 conserved miRNAs in B. mori (Table 3). The relative frequencies of miRNAs generally represent a measure of their expression levels [37,41-43]. We normalized read numbers for all of the miRNAs and compared them on the basis of developmental stages. miRNAs are grouped into following three major categories; The first one is the largest category and with the most dramatic changes in miRNA levels. It includes miR-1, the entire family of miR-2, the miR-9 family (miR-9 and miR-9b), the let-7 family (let-7a, let-7j), miR-10b, miR-31, miR-71, miR-79, miR-87, miR-98, miR-100, miR-252, miR-263a, miR-275, miR-279, miR-317 and miR-1274b (Table 3 and Figure 4a). These miR-NAs are at their lowest levels in larvae but greatly increased in abundance during the spinning larval stage and then decreased their levels during the pupal stage but again sharply increased during the adult stage. The second group consists of miRNAs whose abundance in one of the developmental stages is at least twice high as those of the other three. For instance, miR-276a, miR-274b and miR-923 were preferentially expressed in feeding larvae; miR-1, bantam, miR-8, miR-12, miR-98, miR-190, miR-276, miR-305, miR-970 and miR-iab-4 in spinning larvae; miR-9c, miR-14, miR34, miR-92, miR-277, miR-278 and miR-989 in pupae; and miR-31, miR-100, miR-184, miR-263a and miR-275 in adults (Table 3 and Figure 4b). Notably, miR-8 and miR-1 abundance was greater by 10-fold and 6-fold, respectively, in spinning larvae than in other stages. The levels of several miR-NAs, such as miR-278 (23-fold), miR-981-1 (7 fold) and

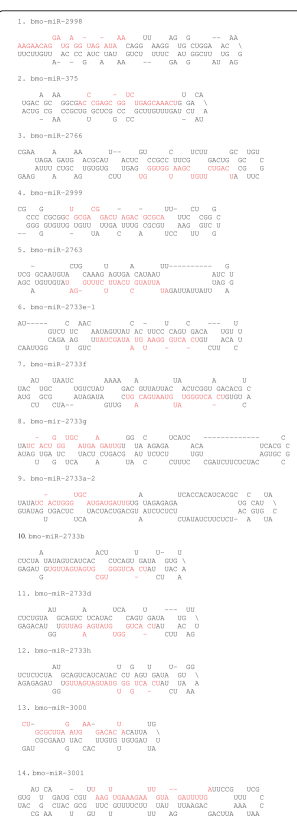


Figure 2 Predicted fold-back structures for the newly identified miRNAs in silkworm. Mature miRNA sequence is shown in red letters.

miR-9c (5-fold), were greater in pupae relative to other stages. miR-100 (5 fold), miR-31 (3 fold) and miR-184 (3 fold) were expressed at much higher levels in moth relative to other stages. Extreme cases of preferential expression were also observed in the dataset. For instance, miR-274a (11 reads) appeared only in feeding larvae, miR-932 (17 reads) only in spinning larvae, miR-210 (7 reads) and miR-449b (8 reads) only in pupae and miR-1000 (4 reads) and miR-206 (2 reads) only in moths (Table 3). Other types of expression patterns also exist: miR-923 level was moderate in feeding larvae, but decreased in spinning larvae maintained at similar levels in pupae and increased in moths (Figure 4c). While miR-116 level steadily increases during progression of larvae to moth (Figure 4d), miR-274b level gradually decreased during the development (Figure 4e), miR-283 and miR-306 levels were relatively uniform in the four stages (Table 3).

miRNAs with extreme low abundance in silkworm libraries

Several of the conserved miRNA, e.g, miR-124, miR-183, miR-193 and miR-210, were expressed at extremely low levels in different stages (Table 3). This could be due to their restricted expression in specific cell-types. For instance, miR-124 and miR-183 were found in retina of mouse and *Drosophila*, respectively [44-46].

Differential expression of let-7 family members during development

An advantage of the sequencing-based approach over hybridization-based methods is that changes in expression profile of individual members within a miRNA family can be determined. Our sequence analysis revealed the expression of at least eight members of the let-7 family (let-7a, -7b, -7c, -7d, -7e, -7f, -7 g and -7j) in *B. mori* (Table 3). Spinning larval and adult stages had the highest numbers of reads for let-7a and let-7j. Reads for let-7b, let-7d and let-7f were recovered only in the moth stage, *albeit* in low numbers.

Expression profile of specific miRNAs differs between *Drosophila*, mosquito and silkworm

MicroRNA, miR-277 expression has been detected only in *Drosophila* adults [20,24]. In contrast, miR-277 is present in the silkmoth at a level much lower than in pupae and spinning larvae (Table 3). In *Anopheles stephensi*, miR-989 is abundantly and specifically expressed in adults [27]. In *B. mori*, the expression of miR-989 was much higher in pupae than in other three stages (Table 3). The miRNA, miR-278 in *Drosophila* is detectable in larvae and increases to peak levels in pupae and adults [47]. In the silkworm, the level of miR-278 was very low in feeding larvae and spinning larvae but increased 23-63 folds in pupae, and then greatly decreased in moths (Table 3). These observations indicate that expression

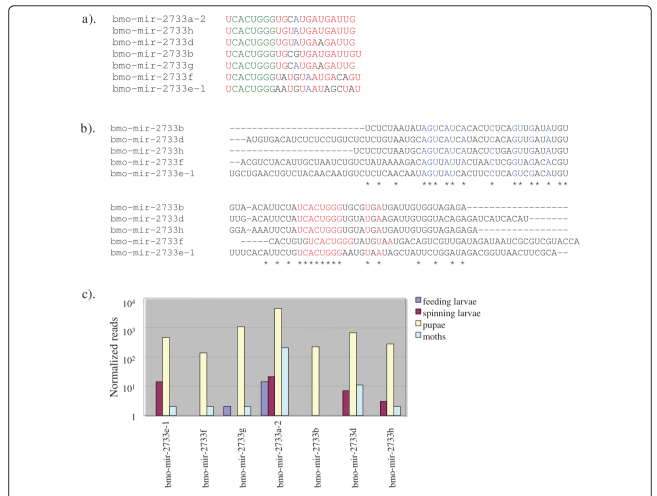


Figure 3 Identification of a novel miRNA family. (a) sequence alignment of the mature miRNAs, (b), sequence alignment of the miRNA precursor (miRNA and miRNA* are shown in red and blue colored letters, respectively) and c), expression levels of the individual members in different developmental stages.

patterns of certain conserved miRNAs differ in insects belonging to different orders taxonomically.

Accumulation of high-level miR* in different developmental stages

Upon processing of a miRNA precursor by Drosha containing complex initially and then by Dicer, the miR: miR* duplex is released, from which only miR is incorporated into RISC. The unincorporated miR* either degrades rapidly or accumulates at significantly low levels [37,38,41,42,48]. Deep sequencing of small RNAs in this study allowed us to monitor the changes in miR* abundance during different developmental stages. Our results largely agree with the previous reports of miR* species recovered at a lower frequency than that of their partners (Table 3 and Figure 5e). However, for several miRNAs i.e, miR-10, miR-33, miR-281 and miR-965, the counts of miR* species far exceeded those of their miR species (Table 3). The relative overall abundance of miR:miR* was in the ratio of 1:48 for miR-33; 1:18 for

miR-281; 1:8 for miR-965 and 1:4 for miR-10a (Figure 5a-d). Intriguingly, high level accumulation is strikingly conspicuous during different developmental stages as shown in Figure 5a-d.

miR-306 and miR-263 loci could generate antisense miRNAs in silkworm

Of the two strands of a miRNA locus, in general, one strand (sense strand) is transcriptionally active. Our sequencing efforts identified, for the first time, the prevalence of at least two new miRNA loci, miR-306 and miR-263, that generate miRNAs by convergent transcription from both strands in the silkworm genome. As well, we recovered eighteen reads of miR-iab-4AS, the first antisense miRNA reported in animals [49]. Analysis of the antisense sequences of miR-306-AS and miR-263-AS loci by mFold showed that they can adopt canonical hairpin secondary structures (Figure 6a and 6b). We recovered eight sequence reads that uniquely mapped to the antisense hairpin sequence of miR-306 (miR-306-

Table 3 Normalized reads of conserved miRNAs during different developmental stages of silkworm (TPM: transcripts per million)

Danitam	miRNA	mature miRNA sequence	Feeding larvae		Spinning larvae		Pupa		Moth	
Demo- et-7a			miR	miR*	miR	miR*	miR	miR*	miR	miR*
Let-7b	bantam	UGAGAUCAUUGUGAAAGCUAAU	168	3	1189	3	210	0	382	6
Let-7c	bmo-let-7a	UGAGGUAGUAGGUUGUAUAGU	817	4	6277	7	748	8	5149	22
Tet-7d	let-7b	UGAGGUAGUAGGUUGUGGUU	0	0	0	0	0	0	2	0
Tet-7e	let-7c	UGAGGUAGUAGGUUGUAUGGUU	7	0	21	0	0	0	63	0
Tet-7f	let-7d	AGAGGUAGUAGGUUGCAUAGUU	0	0	0	0	0	0	4	0
Tet-7g	let-7e	UGAGGUAGUAGGUUGUUUAGUU	2	0	52	0	2	0	17	0
Tet-7 UGAGGUAGUAGGUUGUAUAGUU	let-7f	UGAGGUAGUAGAUUGUAUAGUU	0	0	0	0	0	0	2	0
Tet-7 UGAGGUAGUAGGUUGUAUAGUU	let-7q	UGAGGUAGUAGUUUGUAUAGUU	0	0	17	0	0	0	9	0
miR-1 UGGAAUGUAAAGAAGUUUGAGG 6947 0 80224 3 13618 0 14873 miR-1b UGGAAUGUAAAGAGUAUGUA 1 0 7 0 2 0 0 0 0 0 0 0 0 0 1296 miR-2-1 UAUCACAGCCAGCUUUGAUGAGC 48 1 888 377 127 400 1296 13 0 8 0 127 400 1296 13 0 8 0 127 400 1296 13 0 8 0 291 131 0 8 0 291 131 0 18 0 267 140 146 0 131 0 18 0 267 140 146 0 131 0 18 0 267 145 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	_	UGAGGUAGUAGGUUGUAUAGUU	782	0	6042	0	684	0	4674	0
miR-2-1 UAUCACAGCCAGCUUUGAUGAGC 48 1 888 0 127 0 1296 miR-2-2 UAUCACAGCCAGCUUUGAUGACC 48 20 888 377 127 400 1296 13 miR-2b UAUCACAGCCAGCUUUGAUGAGC 16 0 131 0 8 0 291 13 miR-2d UAUCACAGCCAGCUUUGUUGAUG 9 0 131 0 18 0 267 miR-7 UGGAAGACUAGUGAUUUUGUU 294 1 145 0 75 0 405 miR-7b UGGAAGACUAGUGAUUUUUGUU 294 1 145 0 75 0 405 miR-7b UGGAAGACUAGUGAUUUUUGUU 294 1 145 0 75 0 405 miR-7b UGUUGGUAGUACCAGCUGUAGA 265 9 3342 245 946 25 5125 81 38 3679 456 26 25 5125 81 38 3679 456 26 26 248 297	=	UGGAAUGUAAAGAAGUAUGGAG	6947	0	80224	3	13618	0	14873	0
miR-2-1 UAUCACAGCCAGCUUUGAUGAGC 48 1 888 0 127 0 1296 1 miR-2-2 UAUCACAGCCAGCUUUGAUGAGC 48 20 888 377 127 400 1296 13 miR-2c UAUCACAGCCAGCUUUGAUGGGC 16 0 131 0 18 0 291 miR-2d UAUCACAGCCAGCUUUGUUGUUGUU 9 0 131 0 18 0 267 miR-7b UGGAAGACUAGUGAUUUUGUU 294 1 145 0 75 0 405 miR-7b UGGAAGACUAGUGAUUUUUGUU 2 0 0 0 0 4 4 miR-7b UGGAAGACUAGUAGUAGC 2488 89 35997 321 3738 901 4513 4 miR-9b UCUUUGGUACCUACCUGACUGAGA 265 9 3142 245 946 25 5125 8 miR-9c UCUUUGGUACCUACCUGAGUAGA 10 0 0 0 0	miR-1b	UGGAAUGUUAAGAAGUAUGUA	1	0	7	0	2	0	0	0
miR-2b UAUCACAGCCAGCUUUGAGGAGC 16 0 131 0 8 0 291 miR-2c UAUCACAGCCAGCUUUGAUGGGC 17 0 176 0 12 0 336 miR-2d miR-7d UGGAAGACUAGUGGAUUUUGUUGU 294 1 145 0 75 0 405 miR-7b UGGAAGACUGGGGAAGAGUUGCU 2488 89 35997 321 3738 901 4513 4 miR-9b UCUUUGGUUACUAGCUGUAGA 265 9 3142 245 946 25 5125 5125 8 miR-9b UCUUUGGUUACCUAGCUGUAGA 265 9 3142 245 946 25 5125 58 miR-9b UCUUUGGUUACCUAGCUGAA 195 0 2357 0 539 0 3879 9 1879 0 44 6 1456 76 2334 154 457 6 6 148 77 766 27 1123 3 14 <td>miR-2-1</td> <td></td> <td>48</td> <td>1</td> <td>888</td> <td>0</td> <td>127</td> <td>0</td> <td>1296</td> <td>0</td>	miR-2-1		48	1	888	0	127	0	1296	0
mir-2c UAUCACAGCCAGCUUUGAUGAGU 17 0 176 0 12 0 336 mir-2d UAUCACAGCCAGCUUUGUUGAGU 9 0 131 0 18 0 267 mir-7 UGGAAGACUAGUGAUUUUGUUGU 294 1 145 0 75 0 405 mir-7 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 405 mir-7 UGGAAGACUAGUGAUUUGUUGUU 2 0 0 0 0 0 0 0 4151 44 4451 44 4131 44 4513 44 4513 44 4513 44 4513 44 4513 44 4513 44 4513 44 4513 44 451 46 26 5125 81 4513 44 451 451 451 451 451 451 451 451 451 451 451 451 451	miR-2-2	UAUCACAGCCAGCUUUGAUGAGC	48	20	888	377	127	400	1296	1309
miR-2c UAUCACAGCCAGCUUUGAUGAGG 17 0 176 0 12 0 336 miR-2d miR-2d UAUCACAGCCAGCUUUGUUGAGU 9 0 131 0 18 0 267 miR-7 MAGACUAGUGAUUUUUGUUGU 294 1 145 0 75 0 405 1 miR-7 UGGAAGACUAGUGAUUUUGUUU 2 0 0 0 0 0 0 405 1 miR-70 UGUUGGUACCUAGUAGAAGAUUGU 2488 89 35997 321 3738 901 4513 4 4 miR-9 UCUUUGGUALUCAGCUGUAGA 195 0 2357 0 539 0 3879 miR-10 0 0 0 0 2 0 46 25 5125 8 miR-10 UACCCUUUGUGUAGUUGUAGA 0	miR-2b	UAUCACAGCCAGCUUUGAGGAGC	16	0	131	0	8	0	291	0
miR-2d UAUCACAGCCAGCUUUGUUGAGU 9 0 131 0 18 0 267 miR-7 UGGAAGACUAGUGAUUUUGUUU 294 1 145 0 75 0 405 miR-7b UGGAAGACUAGUGAUUUUUGUU 2 0 0 0 0 0 4 miR-9 UCUUUGGUUACUAGCUGAAAGAUGUC 2488 89 355997 321 3738 901 4513 4 miR-9b UCUUUGGUUACCUAGCUGUAGA 195 0 2357 0 599 0 3879 0 miR-9d UCUUUGGUAUCCUAGCUGUAG 195 0 2357 0 599 0 3879 0 miR-9d UCUUUGGUACCUAGCUAGUUGU 103 27 456 76 2334 154 457 6 miR-10 UACCCUGUAGACCAUACUUGU 123 4 992 7 746 27 1123 3 mir-1b CACCCUGUAGAGUUCUAGCUG 0 0 0 0 <td>miR-2c</td> <td>UAUCACAGCCAGCUUUGAUGGGC</td> <td>17</td> <td>0</td> <td></td> <td>0</td> <td>12</td> <td>0</td> <td>336</td> <td>0</td>	miR-2c	UAUCACAGCCAGCUUUGAUGGGC	17	0		0	12	0	336	0
mir-7 UGGAAGACUAGUGAUUUUGUUGU 294 1 145 0 75 0 405 mir-7b mir-7b UGGAAGACUAGUGAUUUUUGUU 2 0 0 0 0 4 4 mir-8 UAAUACUGUCAGGUGAAGUGUC 2488 89 35997 321 3738 901 4513 4 mir-9b UCUUUGGUACCUAGCUGUAGA 265 9 3142 245 946 25 5125 8° mir-9b UCUUUGGUAUCCUAGCUGUAGA 195 0 2357 0 539 0 3879 mir-9d 0 0 0 0 0 2 0 4 4 457 6 283 154 457 6 mir-9d UCUUUGGUACCUGUUGUG 0 2 1	miR-2d		9	0	131	0	18	0	267	0
mir-7b UGGAAGACUAGUGAUUUUUGUU 2 0 0 0 0 4 mir-8 mir-8 UJAUACUGUCAGGUAAAGAUUC 2488 89 35997 321 3738 901 4513 4 mir-9b UCUUUGGUAUCUAGCUGUAUGA 195 0 2357 0 539 0 3879 0 mir-9c UCUUUGGUAUCUAGCUGUAG 195 0 0 0 0 0 2 0 4 mir-9d UCUUUGGUAUCCUAGCUGUAG 103 27 456 76 2334 154 457 6 mir-10b CACCCUGUCAGAUCUUGUU 60 248 297 726 389 2047 265 9 mir-10b CACCCUGUAGACCUUGUUUUU 123 4 992 7 746 27 1123 1 mir-11b CAUCACAGUCAGAGUUUGUGUU 123 4 992 7 746 27 1123 1 mir-11b CAUCACAGUCAGUUCAGUUGUU 11				1		0		0		6
mir-8 UAAUACUGUCAGGUAAGAUGUC 2488 89 35997 321 3738 901 4513 4 mir-9 UCUUUGGUUAUCUAGCUGUAUGA 265 9 3142 245 946 25 5125 8' mir-9b UCUUUGGUUACCUAGCUGUAGA 195 0 2357 0 539 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 3879 0 457 467 2334 154 457 6 6 2334 154 457 6 9 mir-10 UACCCUGUAGACCUGUGU 60 248 297 7 746 27 1123 1 3 41 7 87 3 123 4 912 7 746 27 1123 3 4 <				0		0		0		0
mir-9 UCUUUGGUUAUCUAGCUGUAUGA 265 9 3142 245 946 25 5125 8 mir-9b UCUUUGGUACCUAGCUGUAGA 195 0 2357 0 539 0 3879 0 mir-9c UCUUUGGUAUCCUAGCUGUAGA 0 0 0 0 2 0 4 457 6 mir-9d UCUUUGGUAUCCUAGCUGAG 103 27 456 76 2334 154 457 6 mir-10b CACCCUGUCAGACCAUCUUGU 60 248 297 726 389 2047 265 9 mir-10b CACCCUGUCAGAGUUCUAGCUA 11 3 4 992 7 746 27 1123 3 mir-11b CAUCACAGUCAGAGUUCUAGCUA 11 3 41 7 87 3 123 mir-13a UAUCACAGCCACUUUGAGUGG 3 0 14 0 10 2 4 1 mir-13a UAUCACAGCCAUUUUUGACGAG				89	35997	321	3738	901		440
miR-9b UCUUUGGUUACCUAGCUGUAUGA 195 0 2357 0 539 0 3879 miR-9c UCUUUGGUAUUCUAGCUGUAGA 0 0 0 0 2 0 4 miR-9d UCUUUGGUAUCCUAGCUGUAG 103 27 456 76 2334 154 457 6 mir-10b CACCCUGUCAGACCUACUUGUU 60 248 297 726 389 2047 265 9 mir-10b CACCCUGUCAGACCUUUGU 123 4 992 7 746 27 1123 3 mir-11 CAUCACAGUCAGAGUUCUGACUA 11 3 41 7 87 3 123 4 mir-11b CAUCACAGCACUUUGAGUUGU 163 0 453 0 125 0 39 1 mir-12 UGAGUAUUACUACAGUCUGUUGUU 163 0 453 0 125 0 39 1 mir-13b UAUCACAGCCACUUUUUGACGGG 28 0 90		UCUUUGGUUAUCUAGCUGUAUGA		9						878
min-9c UCUUUGGUAUUCUAGCUGUAGA 0 0 0 2 0 4 min-9d min-9d UCUUUGGUAUCCUAGCUGUAG 103 27 456 76 2334 154 457 6 min-10 UACCCUGUAGAUCCGAAUUUGU 60 248 297 726 389 2047 265 9 min-10b CACCCUGUCAGACCAUACUUGU 123 4 992 7 746 27 1123 3 min-11b CAUCACAGUCAGAGUUCUAGCUA 0 0 0 0 0 0 0 2 1123 3 11 3 41 7 87 3 123 4 992 7 746 27 1123 3 1123 4 992 7 746 27 1123 3 1 1 3 41 1 7 87 3 123 4 4 4 4 4 4 4 4 4 4 4		UCUUUGGUUACCUAGCUGUAUGA		0		0				0
miR-9d UCUUUGGUAUCCUAGCUGUAG 103 27 456 76 2334 154 457 6 miR-10 UACCCUGUAGAUCCUAGUUGU 60 248 297 726 389 2047 265 9 mir-10b CACCCUGUCAGACCAUACUUGUU 123 4 992 7 746 27 1123 3 miR-11b CAUCACAGUCAGAGUUCUAGCUA 11 3 41 7 87 3 123 3 miR-12b UGAGUAUUACUUCAGGUACUGGU 163 0 453 0 125 0 39 3 miR-13a UAUCACAGCCACUUUGAUGG 3 0 14 0 10 2 4 4 miR-13b UAUCACAGCCACUUUGACUGCUCUCU 100 1 169 221 651 20 39 3 miR-14b UCAGUCUUUUUCUCUCUCUCUA 100 1 169 221 651 20 39 3 miR-31 GGCAGGAGAUGGAUGCGUUGU				0		0		0		0
miR-10 UACCCUGUAGAUCCGAAUUUGU 60 248 297 726 389 2047 265 9 mir-10b CACCCUGUCAGACCAUACUUGUU 123 4 992 7 746 27 1123 3 miR-11 CAUCACAGUCUGAGUUCUAGCUA 0 2 0 0 0 0 0 2 4 0 10 0 14 0 10 0 14 0 10 0 0 14 0 10 0 0 4 13 347 1 0 14 0 10 0 34 <td></td> <td></td> <td></td> <td>27</td> <td></td> <td>76</td> <td></td> <td>154</td> <td></td> <td>67</td>				27		76		154		67
mir-10b CACCCUGUCAGACCAUACUUGUU 123 4 992 7 746 27 1123 3 miR-11 CAUCACAGUCUGAGUUCUUGC 0 0 0 0 0 0 0 0 2 4 miR-11b CAUCACAGUCUGAGAGUUCUAGCUA 11 3 41 7 87 3 123 3 miR-12 UGAGUAUUACUUCAGGUAGUUGGAG 163 0 453 0 125 0 39 miR-13a UAUCACAGCCAUUUUUUGACGAG 28 0 90 0 42 13 347 1 miR-14 UCAGUCUUUUUUCUCUCUCUA 100 1 169 221 651 20 39 3 miR-31 GGCAAGAAGUCGCAUAGCUG 36 0 1801 0 634 0 5617 3 miR-33 GUGCAUGUGUGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-71 UGAAGCUAGGUUGAGAU 2	miR-10		60	248	297	726	389	2047	265	931
miR-11 CAUCACAGUCUGAGUUCUGC 0 0 0 0 0 0 2 description miR-11b CAUCACAGUCAGAGUUCUAGCUA 11 3 41 7 87 3 123 4 miR-12 UGAGUAUUACUUCAGGUACUGGU 163 0 453 0 125 0 39 0 miR-13a UAUCACAGCCACUUUGAUGGG 3 0 14 0 10 2 4 0 miR-13b UAUCACAGCCACUUUUCCCCUA 100 1 169 221 651 20 39 3 miR-14 UCAGUCUUUUCUCUCCCUA 100 1 169 221 651 20 39 3 miR-31 GGCAAGAAGUGGCAUAGCUG 36 0 1801 0 634 0 5617 3 miR-33 GUGCAUUGUGAGUUGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAGCUGGUGAGGUGGUGGUGUU 2 3	mir-10b		123	4	992	7	746	27	1123	2
miR-11b CAUCACAGUCAGAGUUCUAGCUA 11 3 41 7 87 3 123 miR-12 UGAGUAUUACUUCAGGUACUGGU 163 0 453 0 125 0 39 miR-13a UAUCACAGCCACUUUGAUGUG 3 0 14 0 10 2 4 miR-13b UAUCACAGCCAUUUUGACGAG 28 0 90 0 42 13 347 1 miR-14 UCAGUCUUUUUCCUCCUA 100 1 169 221 651 20 39 3 miR-31 GGCAGAGAGUUGCAUGCU 36 0 1801 0 634 0 5617 3 miR-33 GUGCAUUGUAGUUGACUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUGGUUGACUUGCUGUU 106 0 35 0 3981 8 22 1 miR-79 UAAGCCAGUGGUUGUGAGAU 2 3 149 62 48				0		0				0
miR-12 UGAGUAUUACUUCAGGUACUGGU 163 0 453 0 125 0 39 miR-13a UAUCACAGCCACUUUGAUGUG 3 0 14 0 10 2 4 miR-13b UAUCACAGCCAUUUUUGACGAG 28 0 90 0 42 13 347 1 miR-14 UCAGUCUUUUUCUCUCUCCUA 100 1 169 221 651 20 39 miR-31 GGCAAGAAGUCGGCAUAGCUG 36 0 1801 0 634 0 5617 3 miR-33 GUGCAUUGUAGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUUGUAGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAGCUAGUUCCAGGUUGU 2 3 149 62 48 60 86 2 miR-87 GUGAGCAAACUUCAGGUUGU 71 2 131 24 35				3		7	87	3		4
miR-13a UAUCACAGCCACUUUGAUGUG 3 0 14 0 10 2 4 miR-13b UAUCACAGCCAUUUUUGACGAG 28 0 90 0 42 13 347 1 miR-14 UCAGUCUUUUUCUCUCUCUA 100 1 169 221 651 20 39 3 miR-31 GGCAAGAAGUCGGCAUAGCUG 36 0 1801 0 634 0 5617 2 miR-33 GUGCAUUGUAGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUGGUAGCUGGUGU 106 0 35 0 3981 8 22 1 miR-71 UGAAGACAUGAGUUCACAAAGCAU 71 2 131 24 35 7 75 3 miR-79 UAAAGCCAAACUUUCAGGUCUGU 14 0 200 3 52 18 181 3 miR-97 UAUUGCACCAGUCCCGGCCUA 5 0 38 3						0				0
miR-13b UAUCACAGCCAUUUUUGACGAG 28 0 90 0 42 13 347 1 miR-14 UCAGUCUUUUUCCUCUCUA 100 1 169 221 651 20 39 3 miR-31 GGCAAGAAGUCGGCAUAGCUG 36 0 1801 0 634 0 5617 3 miR-33 GUGCAUUGUAGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUGGUAGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAGACAUGGGUAGGUA 2 3 149 62 48 60 86 2 miR-79 UAAAGCUAGAUUCCAAAGCAU 71 2 131 24 35 7 75 3 miR-87 GUGAGCAAACUUUCAGGUGUU 14 0 200 3 52 18 181 3 mir-92 UAUUGCACCAGUCCGGCCUA 5 0				0		0		2		0
miR-14 UCAGUCUUUUUCUCUCUCUA 100 1 169 221 651 20 39 1 miR-31 GGCAAGAAGUCGGCAUAGCUG 36 0 1801 0 634 0 5617 3 miR-33 GUGCAUUGUAGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUGUGUUAGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAAGACAUGGGUAGGAU 2 3 149 62 48 60 86 2 miR-79 UAAAGCUAGAUACCAAAGCAU 71 2 131 24 35 7 75 3 miR-87 GUGAGCAAACUUUCAGGUGUU 14 0 200 3 52 18 181 3 miR-92 UAUUGCACCAGUCCGGCCU 3 0 14 21 33 0 58 4 miR-98 UGAGGUAGAAGUCCGACA 45 0 3 </td <td></td> <td></td> <td></td> <td>0</td> <td></td> <td>0</td> <td></td> <td></td> <td></td> <td>19</td>				0		0				19
miR-31 GGCAAGAAGUCGGCAUAGCUG 36 0 1801 0 634 0 5617 miR-33 GUGCAUUGUAGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUGUGGUUAGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAAGACAUGGGGAGAU 2 3 149 62 48 60 86 2 miR-79 UAAAGCUAGAUUACAAAGCAU 71 2 131 24 35 7 75 3 miR-87 GUGAGCAAACUUUCAGGUUGU 14 0 200 3 52 18 181 3 miR-92 UAUUGCACCAAUCCCGGCCUA 5 0 38 3 120 2 88 miR-98 UGAGGUAGUUGUAUUGUU 16 0 107 0 17 0 67 miR-99 AACCCGUAGAUCCGACCA 45 0 3 0 0 0	miR-14	UCAGUCUUUUUCUCUCCUA		1	169	221	651	20	39	9
miR-33 GUGCAUUGUAGUUGCAUUGC 3 14 17 1013 3 43 0 2 miR-34 UGGCAGUGUGGUUAGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAAGCAUGGGUAGUGAGAU 2 3 149 62 48 60 86 2 miR-79 UAAAGCUAGAUUCCAAACUU 71 2 131 24 35 7 75 5 miR-87 GUGAGCAAACUUCAGGUGUGU 14 0 200 3 52 18 181 3 miR-92 UAUUGCACCAGUCCCGGCCUA 5 0 38 3 120 2 88 miR-92 AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 miR-98 UGAGGUUGAAAGUCGCACA 45 0 3 0 3 0 4 miR-99 AACCCGUAGAUCCGAGCUGUU 0 0 3 0 0 0				0						2
miR-34 UGGCAGUGUGGUUAGCUGGUUG 106 0 35 0 3981 8 22 1 miR-71 UGAAAGACAUGGGUAGUGAGAU 2 3 149 62 48 60 86 2 miR-79 UAAAGCUAGAUUACCAAAGCAU 71 2 131 24 35 7 75 3 miR-87 GUGAGCAAACUUCCAGUGUGU 14 0 200 3 52 18 181 3 miR-92 UAUUGCACCAGUCCCGGCCUA 5 0 38 3 120 2 88 6 mir-92b AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 6 miR-98 UGAGGUAGAGUUGUAUUGUU 16 0 107 0 17 0 67 67 miR-98 family UGAGGUAGAGUCCGACCA 45 0 3 0 0 0 11 6 miR-190 AACCCGUAGAUCCGACUUGU 0 0 3<				14						26
miR-71 UGAAAGACAUGGGUAGUGAGAU 2 3 149 62 48 60 86 2 miR-79 UAAAGCUAGAUUACCAAAGCAU 71 2 131 24 35 7 75 3 miR-87 GUGAGCAAACUUUCAGGUGUGU 14 0 200 3 52 18 181 3 miR-92 UAUUGCACCAGUCCCGGCCU 3 0 38 3 120 2 88 6 mir-92b AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 6 miR-98 UGAGGUAGGUUGUAUUGUU 16 0 107 0 17 0 67 67 miR-98 family UGAGGUUGAAAGUCGACAC 45 0 3 0 3 0 4 67 </td <td>miR-34</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>11</td>	miR-34									11
miR-79 UAAAGCUAGAUUACCAAAGCAU 71 2 131 24 35 7 75 miR-87 GUGAGCAAACUUUCAGGUGUGU 14 0 200 3 52 18 181 miR-92 UAUUGCACCAGUCCCGGCCU 5 0 38 3 120 2 88 mir-92b AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 miR-98 UGAGGUUGAUGUGUUUUU 16 0 107 0 17 0 67 miR-98 family UGAGGUUGAAAGUCGACA 45 0 3 0 3 0 4 miR-99 AACCCGUAGAUCCGAGCUUGUU 0 3 0 0 0 11 miR-100 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 miR-133 UAUUGCUUGAGAAUACACGUG <th< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td>22</td></th<>										22
miR-87 GUGAGCAAACUUUCAGGUGUGU 14 0 200 3 52 18 181 miR-92 UAUUGCACCAGUCCCGGCCUA 5 0 38 3 120 2 88 mir-92b AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 miR-98 UGAGGUAGGUGUAUUGUU 16 0 107 0 17 0 67 miR-98 family UGAGGUUGAAAGUCGACA 45 0 3 0 3 0 4 miR-99 AACCCGUAGAUCCGAGCUUGUU 0 0 3 0 0 0 11 miR-100 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-183 UAUGGCACGGAGAAUGACA										9
mir-92 UAUUGCACCAGUCCCGGCCUA 5 0 38 3 120 2 88 mir-92b AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 mir-98 UGAGGUAGGUAGGUUGUAUUGUU 16 0 107 0 17 0 67 mir-98 family UGAGGUUGAAAGUCGCACA 45 0 3 0 3 0 4 mir-99 AACCCGUAGAUCCGAGCUUGUU 0 3 0 0 0 11 mir-100 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 mir-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 mir-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 mir-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 mir-183 UAUGGCACUGGUAGAAUUCACU <										2
mir-92b AAUUGCACCAAUCCCGGCCU 3 0 14 21 33 0 58 miR-98 UGAGGUAGUAGGUUGUAUUGUU 16 0 107 0 17 0 67 miR-98 family UGAGGUUGAAAGUCGACA 45 0 3 0 3 0 4 miR-99 AACCCGUAGAUCCGAACUUGUU 0 0 3 0 0 0 11 miR-100 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC										0
miR-98 UGAGGUAGGUUGUAUUGUU 16 0 107 0 17 0 67 miR-98 family UGAGGUUGAAAGUCGCACA 45 0 3 0 3 0 4 miR-99 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-100 AACCCGUAGAUGCCAA 1 0 3 0 1 0 0 0 1 0 1 0 7 0 3 0 1 0 0 0 0 1 1 0										4
miR-98 family UGAGGUUGAAAGUCGCACA 45 0 3 0 3 0 4 miR-99 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-99 AACCCGUAGAUCCGAGCUUGUU 0 0 3 0 0 0 11 miR-100 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-100 AACCCGUAGAUCCGAACUUGUG 128 0 636 3 405 0 3223 miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 0 miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507	-									0
miR-124 UAAGGCACGCGGUGAAUGCCAA 1 0 3 0 0 0 0 miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-133 UUUGGUCCCCUUCAACCAGCUG 15 1 55 0 15 0 11 miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-137 UAUUGCUUGAGAAUACACGUAG 1 0 7 0 3 0 11 miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-183 UAUGGCACUGGUAGAAUUCACU 3 0 0 0 5 0 41 miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-184 UGGACGGAGAACUGAUAAGGGC 611 0 3743 3 2660 5 11507										0
miR-190 AGAUAUGUUUGAUAUCUUGGUUG 22 8 197 55 47 100 34										0 9

Table 3: Normalized reads of conserved miRNAs during different developmental stages of silkworm (TPM: transcripts per million) (Continued)

miR-206 UGGAAUGUAAGGAAGUGUGUGG 0 0 0 0 0 2	0 0
miR-210 UGUGCGUGUGACAGCGGCUA 0 0 0 7 0 0	0
mir-228 AAUGGCACUGCAUGAAUUCACGG 1 0 17 0 8 0 28	0
miR-236 UAAUACUGUCAGGUAAUGACGCU 3 0 24 0 7 0 2	0
miR-252 CUAAGUACUAGUGCCGCAGGAG 53 0 463 0 224 0 802	0
miR-263a AAUGGCACUGGAAGAAUUCAC 1177 2 7376 0 2698 0 22066	2
miR-263b CUUGGCACUGGGAGAAUUCAC 6 1 31 17 108 10 65 1	.7
miR-274 UUUUGUGACCGACACUAACGGGUAAU 11 0 0 0 0 0 0	0
miR-274b UUUGUGACCGUCACUAACGGGCA 823 29 83 45 5 0 0	0
miR-275 UCAGGUACCUGAAGUAGCGCGCG 32 1 1856 17 215 0 2083	2
miR-276 AGCGAGGUAUAGAGUUCCUACG 26 27 73 73 58 58 15 1	.7
miR-276a UAGGAACUUCAUACCGUGCUCU 2108 0 30401 0 22507 0 19446	0
miR-276b UAGGAACUUAAUACCGUGCUCU 1 0 21 0 25 0 22	0
miR-277 UAAAUGCACUAUCUGGUACGACA 111 0 3336 7 6488 47 1815	2
miR-278 UCGGUGGGAUCUUCGUCCGUU 105 78 283 1421 6648 219 37 8	60
miR-278 UCGGUGGGAUCUUCGUCCGUUU 103 71 280 1269 6546 179 34 7	37
miR-278 UCGGUGGGAUUUUCGUCCGUUU 2 0 3 0 159 0 2	0
miR-279 UGACUAGAUCCACACUCAU 16 0 408 3 204 5 608	6
miR-279 family UGACUAGAUUUUCACUUAUCCU 14 12 33 114 15 25 29 6	3
miR-281 CUGUCAUGGAGUUGCUCUUUUA 12 303 17 518 65 916 11 1	88
miR-282 ACCUAGCCUCCUUGGCUUUGUCUGU 192 4 577 7 154 2 60	0
miR-283 UAAAUAUCAGCUGGUAAUUCUGGG 222 2 401 0 169 0 237	0
miR-285 UAGCACCAUUCGAAUUCAGUG 1 0 0 0 10 0 0	0
miR-286 GGACUGGAUCCGGACCCGCGUUCUC 1743 0 332 0 200 0 82	0
miR-305 AUUGUACUUCAUCAGGUGCUCUG 39 1 1863 62 28 2 302 4	5
miR-306 UCAGGUACUAGGUGACUCUGA 2542 3 3256 10 2662 8 4401	4
miR-307 UCACAACCUCCUUGAGUGAG 50 0 944 7 479 0 1332	2
miR-308 AAUCACAGGAUAAUACUGCGAG 134 6 156 31 47 28 50 2	22
miR-309 UCACUGGGUGCAUGAUGAUCGU 0 0 0 327 0 0	0
miR-317 UGAACACAGCUGGUGGUAUCC 117 0 1358 0 671 0 1367	0
miR-429 UAAUACUGUCUGGUAAUGCCG 0 0 3 0 0 0	0
miR-449b AGGCAGUGUUGUUAGCUGGC 0 0 0 0 8 0 0	0
miR-923 GUAAGCGGAGGAAAAGAACU 1214 0 294 0 255 0 839	0
miR-927 UUUAGAAUUCCUACGCUUUACC 29 30 131 73 75 50 75 5	8
miR-932 UCAAUUCCGUAGUGCAUUGCAG 0 0 17 0 0 0	0
miR-965 UAAGCGUAUAGCUUUUCCCCUU 16 14 21 283 10 135 6 6	0
	0
miR-980 CAGCUGCCUAGCGAAGGGCAA 54 47 20 69 14 23 17 3	37
miR-981 UUCGUUGUCGAAACCUGCAA 2 0 10 0 30 0 17	0
miR-989-1 UGUGAUGUGACGUAGUGGAA 2 0 477 0 3494 2 45	0
miR-989-2 UGUGAUGUGACGUAGUGGAA 2 0 477 0 3494 0 45	0
miR-993-1 GAAGCUCGUCUCUACAGGUAUCU 0 19 7 169 8 100 6 1	64
miR-993-2 AAAGCUCGUCUCUACAGGUAUA 0 19 7 176 8 100 6 1	81
miR-998 UAGCACCAUGGGAUUCAGCUC 12 0 28 0 110 0 54	0
miR-1000 AUAUUGUCCUGUCACAGCAGU 0 0 0 0 0 4	0
miR-1175 UGAGAUUCAACUCCUCCAACUUAA 28 189 14 14 2 0 9	0
	0
	0
miR-iab-4 ACGUAUACUGAAUGUAUCCUGA 17 1 359 3 42 0 6	0
miR-iab-4-3p CGGUAUACCUUCAGUAUACGUAAC 1 17 3 356 0 43 0	6
miR-iab-4as-5p UUACGUAUACUGAAGGUAUACCG 1 0 17 0 0 0	0

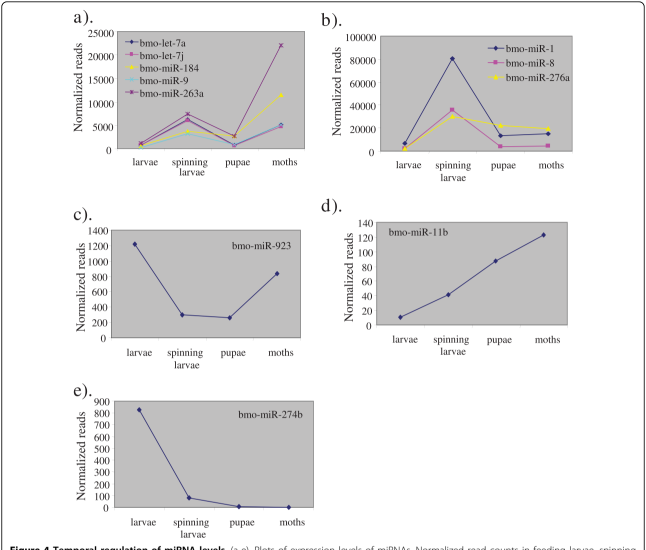


Figure 4 Temporal regulation of miRNA levels. (a-e), Plots of expression levels of miRNAs. Normalized read counts in feeding larvae, spinning larvae, pupae and moths are used for the graphical presentation.

AS) and a single read mapping to the antisense hairpin of miR-263 (miR-263-AS) (Figure 6c and 6d), which indicates that the antisense transcripts were processed into mature miRNAs *in vivo*. miR-306 is fairly uniformly expressed across the developmental stages, as judged by the number of reads, whereas miR-263 is dynamically regulated from the larva to adult stages. In both cases, the recovery of reads from the antisense strand was much lower than that of the sense strand.

Comparing the mature sequences in each sense and antisense pair showed the seed region in miR-263 and miR-263-AS to be almost identical, whereas miR-306 and miR-306-AS showed variations in several internal positions, including the seed region (Figure 6e). In *Drosophila*, miR-iab-4-AS is functional in repressing the Hox family of genes [50,51]. miR-iab-4 and miR-iab-4-AS are

not identical in their seed region, so they are predicted to target a different set of genes [51]. The same could be true for miR-306 and miR-306-AS. Identification of three antisense miRNAs for miR-306, miR-263, and miR-iab-4-AS in silkworm suggests that such antisense miRNAs may be more widespread in other organisms.

Are miR-1920 and miR-1921 genuinely miRNAs in silkworm?

Mapping small RNA reads in this study revealed that certain small RNAs previously annotated as miRNAs may not qualify for such an annotation. In a recent study, two small RNAs were annotated as miR-1920 (one read) and miR-1921 (three reads) in silkworm [35]. Our sequences represented not only the annotated miR-1920 and miR-1921 but also several other small RNAs from these loci. Closer inspection of small RNAs

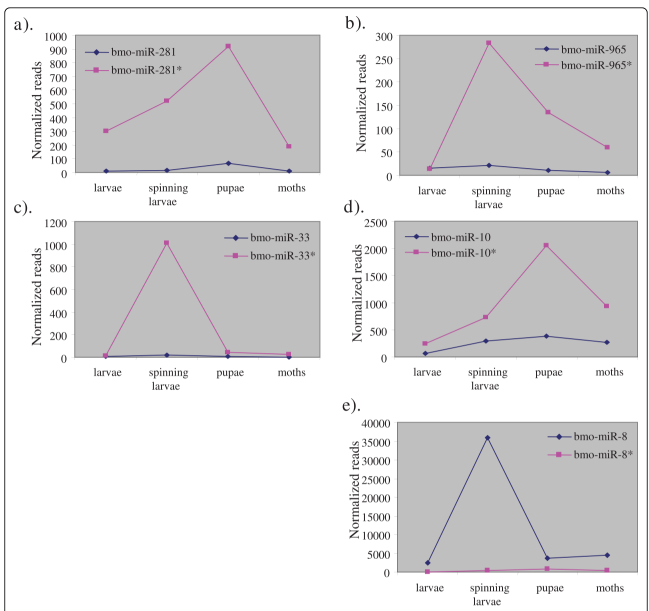


Figure 5 Temporal regulation of miRNA-star (miR*) abundance (a-e), Plots of expression levels of miRNAs along with their miRNA* reads. Normalized read counts in feeding larvae, spinning larvae, pupae and moth stages in silkworm are used for the graphical representation.

generated from these loci revealed that multiple overlapping small RNA reads could be mapped to almost entire length of the hairpin sequence (Figure 7). These results suggest that the annotated miR-1920 and miR-1921 likely arose from siRNAs generating genomic loci or by fortuitous cleavage of transcripts and not strictly from stem-loop biogenesis. Thus, deep sequencing can aid in confident annotation of small RNAs into miRNAs.

Discussion

In order to build a normal animal, development of individual tissues and organs needs to be tightly coordinated

with the developmental progression of the whole organism [52]. This process is even more critical in insect species, including *B. mori*, that undergo well-defined developmental stages (metamorphosis). To gain insight into post-transcriptional gene regulation in silkworm development, determining a near-complete set of miR-NAs and their expression patterns is essential.

Novel miRNAs in silkworm are preferentially expressed in the pupal stage

Recent studies in Drosophilid species led to the suggestion that most novel miRNAs originated from non-miRNA sequences and only a small fraction (4%) is

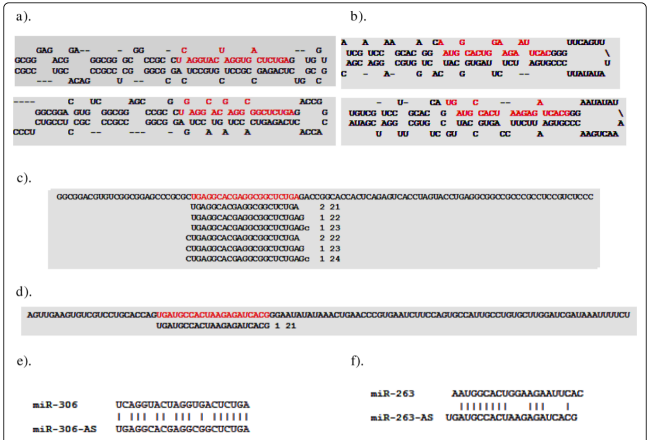


Figure 6 Antisense miRNAs in *B. mori*. Predicted fold-back of miR-306 (a) and miR-263 (b) from sense (top panel) and antisense strand (bottom panel) and mature miRNA is shown in red. Sequencing reads uniquely aligned to hairpin sequence of reverse complement of miR-306 (c) and miR-263 (d) followed by their cloning frequency and length (nt). Mature sequences of miR-306:miR-306-AS pair (e) and miR-263:miR263-AS (f) aligned for showing identical nucleotide sequences.

expected to be retained during evolution [53]. Only 2.5% of the surviving miRNAs are expressed at moderate levels [53]. The observation that most of the newly identified silkworm-specific miRNAs were expressed as abundantly as some of the conserved miRNAs implies that these novel miRNAs evolved, survived, and possibly integrated into the silkworm post-transcriptional regulatory networks. Interestingly, most of these miRNAs showed highest expression in the pupae stage, suggesting a complex gene regulation involving species-specific miRNAs in the development of adult structures.

We have predicted over 1100 genes as targets for the newly identified silkworm-specific miRNAs and their stars using hitsensor software [54] (Additional file 1). Cloning of several novel miRNAs specifically from pupal stage suggests an important role for these miRNAs in larval-pupal-adult metamorphosis. Consistent with this suggestion, many predicted targets for novel miRNAs are genes essential for insect development, molting and metamorphosis, which are regulated by two hormones

(ecdysone and its active metabolite 20-hydroxyecdysone (20E) and the sesquiterpenoid juvenile hormone (JH) [55]. A novel miRNA, bmo-mir-2763 is predicted to target diapause hormone receptor-4. Bombyx diapause hormone receptor (BmDHR) is expressed in the prothoracic gland (PG), the organ, which synthesizes and releases the insect molting hormones, ecdysteroid [56]. JH synthesis and degradation are two routes that control JH levels in the insect haemolymph (Sheng et al., 2008). Juvenile hormone acid O-methyltransferase (JHAMT) is a key enzyme involved in JH synthesis and is a predicted target for miRNA, bmo-miR-2998. The expression profile of JHAMT mRNA in silkworm and Drosophila melanogaster suggest that the suppression of JHAMT transcript is critical for the induction of larval-pupal metamorphosis [57]. Juvenile hormone esterase (JHE) hydrolyses the JH, thus regulates the levels of JH [58]. JHE is a predicted target for bmo-mir-2766 in silkworm. Thus, the genes encoding enzymes implicated in JH biosynthesis and degradation are predicted targets for novel

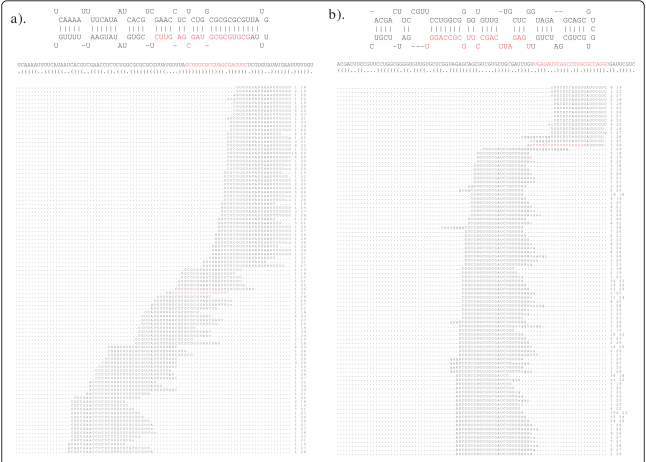


Figure 7 Overlapping small RNA reads derived from annotated miR-1920 and miR-1921 loci. Annotated mature miRNAs are shown in red in the predicted fold-back structures for miR-1920 (a) and miR-1921 (b). Overlapping small RNA reads derived from the predicted hairpin structure are shown in the bottom panel and number of reads recovered in the libraries and their length (nt) are shown on the right.

miRNAs in silkworm. These target predictions suggest that JH levels are regulated by two novel miRNAs that are specifically and abundantly expressed in pupal stage.

A predicted target for novel miRNA (bmo-mir-2763) is nuclear receptor GRF (germ cell nuclear factor (GCNF)-related factor). Increasing evidence suggests that GCNF is important for both female and male reproduction [59]. The developmental program of spermatogenesis is regulated by several transcription factors, one of which appears to be GCNF [59]. GCNF is a sequence-specific repressor of transcription, which binds as a homodimer or an oligomer to its cognate response elements [59]. Identification of GCNF as a target for silkworm-specific miRNA specifically expressed in pupal stage suggests a role for this miRNA in development of male and female reproduction.

During the terminal stages of differentiation of the silkworm ovary, the follicular cells that surround the oocyte produce a large number of related polypeptides that are involved in the formation of the eggshell or chorion [60]. MicroRNA, bmo-mir-2763 is predicted to target a BmGATA beta isoform 3, which regulates the expression of a class of chorion genes expressed during the late stages of choriogenesis [60]. Consistent with the detection of bmo-mir-2763 in the pupal stage, GATAbeta 3 in silkworm has been detected in pupae, but none of the larval tissues [60], suggesting that this process is restricted to pupal stage and novel miRNA (bmo-mir-2763) has a role in the process of choriogenesis. Another miRNA, bmo-mir-2733e-1 is predicted to target NADPH cytochrome P450 reductase, a component of the microsomal P450 electron transport system. It plays an essential role in the transfer of reducing equivalents from NADPH to various P450 molecules and said to be involved on in the ecdysone 20-hydroxylation during the insect's embryonic development [61]

The predicted target for bmo-miR-2998* is silk gland factor-1 (SGF-1). SGF-1 is a member of the fork head/HNF-3 family, which regulates transcription of tissue-specific genes [62]. It plays a role in organogenesis

processes such as those of the gut, silk glands, and nervous systems, act as a region-specific homeotic gene [63]. It was also speculated that SGF-1 protein may be initially required for the development of silk glands and subsequently utilized in the control of genes coding for silk proteins. The miRNA, bmo-miR-2998* can potentially target Argonaute-2. Argonaute (Ago) 2 is the catalytic engine of RNA interference, but little is known concerning the regulation of Ago2 by miRNAs or miRNA* in animals, whereas in plants miR168 can target AGO-1, the major Argonaute involved in target mRNA degradation. Recent studies indicate that Ago2 is required for a variety of developmental processes that occur in a tissue-specific manner [64]. These predictions indicate that several of the new miRNAs and miRNA* are likely to play potentially important roles in metamorphosis by regulating genes that participate in these processes.

Differential regulation of miRNAs during silkworm development implies a role for miRNAs in diverse aspects of *B. mori* physiology and development

Evolutionarily, miR-1 is highly conserved and involved in muscle development and physiology [65,66]. The sole miR-1 gene in *Drosophila* is transcriptionally up-regulated by *Twist* and *Mef2* during myogenesis [66]. In addition to miR-1, miR-133 and miR-206 are designated as muscle-specific miRNAs because of their abundance in diverse muscle tissues [38,67]. In *B. mori*, miR-1 level is very high, miR-133 is low and miR-206 is extremely low, which suggests that miR-1 likely plays a major role in muscle development and physiology in insects.

Holometabolous insects including silkworm undergoes apoptosis during development from larvae to pupae to moth. The spinning larval stage is a stage of transition to pupae during which gut, salivary glands, the silk gland and larval-specific muscles undergo programmed cell death and histolysis [68]. Several miRNAs known to play a role in apoptosis are expressed in a stage-dependent manner in silkworm. miR-2 and miR-13 families possess identical "seed" sequences, and both show almost similar expression profiles in the silkworm. The miR-2 family is known to target proapoptotic genes (reaper, grim and sickle) in Drosophila [69]. Preferential expression of miR-2 and miR-13 families in spinning larvae and moth suggests that these miRNAs may promote apoptosis in these two stages. Another miRNA, miR-34 in C. elegans, plays a role in apoptotic and nonapoptotic cell death [70]. In B. mori, miR-34 is preferentially and abundantly expressed in pupae relative to the other three stages (Table 3), and this expression profile is similar to what was reported for C. elegans [70]. Thus, different miRNAs may play similar roles but in a developmental stage-specific manner in B. mori.

Wg/Wnt signaling is highly regulated and inappropriate activation or inhibition of the pathway results in developmental defects and diseases [71,72]. miR-8 is a negative regulator of Wnt signaling in *Drosophila* [73]. Abundant miR-8 during the silkworm spinning-larval stage suggests a role in Wnt signaling. The seed sequences of miR-8 and miR-429 are identical, indicative of a potential in targeting similar sequences (messenger RNA targets). However, in the silkworm, the two miR-NAs vary greatly in abundance: miR-8 level was high whereas miR-429 level was extremely low. Perhaps, miR-8 plays a predominant role in Wnt signaling in *B. mori*.

In Drosophila, miR-14 and miR-278 are implicated in fat/energy metabolism [37]. miR-278 knockout flies are lean suggesting an excessive utilization of lipid stores. The similarity between the expression profiles of miR-278 in silkworm (Table 3) and Drosophila [37] suggests that miR-278 could play a role in regulating energy metabolism in silkworm. miR-14 is also preferentially expressed in pupae of silkworm, which suggests that miR-14 could also play a role in fat metabolism. On the basis of the roles of these two Drosphila's miRNA, it is tempting to speculate that the increased levels of miR-14 and miR-278 observed in the pupal stage could prevent an excessive utilization of fat. This hypothesis would also be consistent with studies in lepidopteran and other insect classes that have shown a preferential utilization of glycogen stores during pupation (Arrese and Soulages, Annual Rev. Entomology, in press). By preventing the utilization of fat, needed for reproduction during the adult stages, these miRNAs would promote the utilization of glycogen stores.

Of all four developmental stages we analyzed, the spinning-larval stage is important - silk is synthesized and a cocoon is built during this period. Strikingly, the overall miRNA abundance is almost doubled from feeding to spinning larval stages, which suggests a role for miRNAs in silk synthesis, besides their importance in developmental transition. The increase in miRNA levels is mostly quantitative and limited to conserved miRNAs. The levels of novel miRNAs in spinning larvae were very low. Similarly, miRNA levels are also sharply elevated in moths relative to the pupae. Again the increase could not be attributed to qualitative differences. Many conserved miRNAs along with a few species-specific miRNAs become more abundant and may regulate reproduction or aging.

The most abundantly expressed miRNAs in B. mori are also highly conserved miRNAs e.g, miR-1, miR-8, miR-10, let-7, miR-263a, miR276a, and miR-306 and were expressed in all four stages, *albeit* their expression levels vary across different developmental periods. These abundantly expressed miRNAs accounted for most of

the differences in miRNA abundance in different developmental stages of the silkworm. Variation in the level of a specific miRNA across developmental stages indicates stage-specific function, whereas a similar expression over a range of developmental stages points to a possible role in basal and/or constitutive processes. This analysis also revealed that many miRNAs are similarly regulated during development, suggestive of possible coregulation of these miRNAs during silkworm development (Figure 4a and 4b). The changes in miRNA expression profile could imply a fine-tuning or tight regulation of their targets in a spatial- and temporal-specific manner. Taken together, conserved miRNAs along with the species-specific miRNAs, in part, orchestrate the developmental progression of *B. mori*.

miR* levels for several miRNAs are dynamically regulated during development

The specific accumulation of miR* sequences for miR-8, miR-10, miR-37, miR-281 and miR-965 was particularly striking in the silkworm (Table 3 and Figure 5). miR-10* had a higher frequency than miR-10 in Drososphila [23,74] and locust [31]. These observations suggest that miR-10* tends to accumulate at a higher level. This implies a potential functional role for miR-10*, and possibly other miR* that tends to accumulate at significant levels. Indeed, Okamura et al. [74] recently demonstrated a regulatory role for miR-276*, which can potentially target a gene and negatively regulate its expression. For several miRNAs, we found that the accumulation of miR* did not correspond with the miR levels in different stages of *B*. *mori.* miR* accumulation may also be developmentally regulated in the silkworm, which suggests a more complex regulation of miR and miR* sequences than was previously proposed. The observation that several miR*s are more abundant than their corresponding miRNAs indicates that silkworm miR*s could have some endogenous targets. Because of sufficient complementarity between miR and miR* sequences, the accumulation of miR* could potentially regulate the miRNA precursor itself, as was shown for few cases in Arabidopsis [75] and rice (Li and Sunkar, unpublished data).

Comparision between miRNA array and sequencing-based profiling in the silkworm

Recently, miRNA microarray was used to profile their expression during silkworm metamorphosis [33,36]. The expression patterns of several miRNAs differed between the two previous reports and this study [33,36]. For instance, using miRNA array analysis Zhang et al. [36] reported that miR-9c was highly expressed in larvae, although it was also present in eggs and pupae. This observation prompted the authors to suggest that miR-9c is involved in regulation of metamorphosis from larvae to pupae. However, our sequencing-based analysis of miR-9c abundance sharply contrasts with that in the

previous report (Table 3 Figure 4a). First, miR-9 family represented by three members, miR-9, miR-9b and miR-9c in silkworm and these members differ by one nucleotide. Thus, it is difficult to determine which of these members is being detected from a hybridization-based approach. Second, miR-9c expression is extremely low: we recovered one read from the pupae and two reads from the adults, and none from the feeding or spinning larvae (Table 3). Third, we recovered abundant miR-9 and miR-9b reads from all four libraries, and both exhibited a dynamic regulation (Figure 4). Overall, miR-9b level was lower than that of miR-9 in any given stage. miR-9 family had the highest expression in moths and the lowest in larvae. In another miRNA array-based study, He et al., [33] demonstrated that let-7b and let-7d were expressed only in larval and pupal stages but not in the adult stage. Since let-7 family is represented by eight members in the silkworm, each of which differed by one nucleotide, which makes distinguishing difficult by the hybridization-based approach. On the basis of read counts, only let-7a and let-7j are highly abundant, and the remaining six members are low (Table 3). In contrast to what was reported [33], we recovered reads for both let-7b and let-7d from the adult library (albeit in low numbers) but none from the larval or pupal libraries. Another miRNA, miR-277, was reported to be specifically expressed in the moth stage [33]. We detected miR-277 in all four stages. The level was highest in pupae and estimated to be threefold greater than that in the moths. Similarly, the expression profiles of several other miRNAs, as deduced from the array analysis, did not match our analysis. Perhaps, the array-based approach [33,36] needs further validation and we believe that the sequencing-based study is more reliable. Results from miRNA array analysis are likely affected by false signals, cross-hybridizations or background noise.

Conclusions

Deep sequencing of small RNAs from four distinct developmental stages of the silkworm (feeding larvae, spinning larvae, pupae and moths), each with distinct anatomy, morphology and physiology, allowed us to comprehend changes in the miRNA expression profiles during development and to discover novel miRNAs in B. mori. Our study represents the first exhaustive expression profiling of miRNAs in the major developmental stages. We discovered 14 novel miRNAs in the silkworm that are potentially species-specific, identified 101 conserved miRNAs, monitored the dynamic changes in miRNA levels during metamorphosis, and identified miR* species that accumulate at higher levels than their corresponding miRNAs. The accumulation of miR*appears to be regulated in a stage-specific manner. In addition, we found novel antisense miRNA loci in

silkworm. Future identification of the miRNA targets should enable us to integrate the data into post-transcriptional regulatory networks important for silkworm metamorphosis, as well as other economically important traits such as silk synthesis.

Methods

Insect rearing, total RNA preparation, and miRNA library construction

B. mori eggs and diet were purchased from Carolina Biological Supply (Burlington, NC) and larvae were reared at ambient temperature. Whole insects in the feeding larval (4th and 5th instars), spinning larval, pupal, and adult stages were separately frozen in liquid nitrogen and grinded for total RNA extraction using Trizol reagent (Invitrogen Life Technologies, Carlsbad, CA). Total RNA was dissolved in de-ionized formamide for effective RNA denaturation. Small RNAs in the desired size range (15-30 nt) were purified from denaturing 15% polyacrylamide gel and sequentially ligated with the 3' and 5' adapters as described previously [38]. Reverse transcription reaction was performed using the RT primer (5'CAAGCAGAAGACGGCATACGA), and the forward and reverse primers (5'AATGATACGGC-GACCACCGACAGGTTCAGAGTTCTACAGTCCGA and 5'CAAGCAGAAGACGGCATACGA) primers. After phenol/chloroform extraction and ethanol precipitation, the PCR products were shipped to Illumina (city, CA) for sequencing.

Sequence analysis and identification of homologs of conserved miRNAs in *B. mori*

Our computational methods for analyzing the small RNA library was reported previously [38]. In brief, all small RNA reads without perfect matches to the most proximal 11 nt of the 5' adaptor sequences were first removed. Reads corresponding to repeats were discarded using the Einverted and Etandem programs in the EMBOSS package. The unique small RNAs were aligned to RepBase v13.04 [76] and known non-coding RNAs (rRNAs, tRNAs, snRNAs, snoRNAs, etc.) obtained from Rfam [77] with NCBI BLASTN. After removal of repetitive elements, the remaining small RNAs were mapped to the messenger RNAs and such sequences were discarded assuming that these represent degradation products. The filtered sequences were mapped to the reported miRNAs in the miRBase v13. Small RNAs that matched known miRNAs of the silkworm or other animal species resulted in identification of conserved miRNA homologs in *B. mori*.

Identification of novel miRNAs in B. mori

The remaining unique small RNAs were aligned to genome sequences of *B. mori* downloaded from Silkworm Genome Database [78] and for those sequences that matched with the genome, the fold-back structures were

predicted using their flanking sequences [79]. Candidates with predictable fold-back structures were analyzed for the presence of miR* sequences in library. Initially, for such candidates we predicted miR* sequences with 2 nt 3' overhangs on the opposite arm of the fold-back structures and subsequently we searched for the presence of such miR* sequences in our library. This resulted in identification of 12 miRNAs. Additional two small RNAs (bmo-mir-2733e-1 and bmo-mir-2733f) without their miR* were added to the list of new miRNAs, because these two small RNAs are members of the newly identified miRNA family in *B. mori.*

Additional file 1: Predicted targets for the newly identified miRNAs and miRNA*s in silkworm. For each predicted target, accession number, annotation and hit scores are provided.

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[http://www.biomedcentral.com/content/supplementary/1471-2164-11-52-S1.XLS]

Additional file 2: Top 20 most abundantly expressed miRNAs in silkworm during different developmental stages. Proportion of reads recovered for miRNAs in each of the four developmental stages, i.e., feeding larvae, spinning larvae, pupae and moth are provided. Click here for file

[http://www.biomedcentral.com/content/supplementary/1471-2164-11-52-S2.PPT]

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Authors' contributions

R.S. designed the research, analyzed the data, coordinated the project and wrote the paper. G.J. constructed the small RNA library. Y.Z. and W.Z. performed the computational analysis. N.S and H.J. reared the insects and isolated total RNA used for library construction. H.J. also edited the manuscript. J.L.S. and E.A helped with the analysis and interpretation of the computational dataset. All authors read and approved the final manuscript.

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References

- Flynt AS, Lai EC: Biological principles of microRNA-mediated regulation: shared themes amid diversity. Nat Rev Genet 2008, 9:831-842.
- Siomi H, Siomi MC: On the road to reading the RNA-interference code. Nature 2009. 457:396-404.
- Banerjee D, Slack F: Control of developmental timing by small temporal RNAs: a paradigm for RNA-mediated regulation of gene expression. *Bioessays* 2002, 24:119-129.
- Banerjee D, Slack FJ: Temporal and spatial patterning of an organ by a single transcription factor. Genome Biol 2005, 6:205.
- Bartel DP: MicroRNAs: genomics, biogenesis, mechanism, and function. Cell 2004, 116:281-297.

- Bartel DP: MicroRNAs: target recognition and regulatory functions. Cell 2009. 136:215-233.
- Cordes KR, Srivastava D: MicroRNA regulation of cardiovascular development. Circ Res 2009, 104:724-732.
- Hagen JW, Lai EC: microRNA control of cell-cell signaling during development and disease. Cell Cycle 2008, 7:2327-2332.
- Sunkar R, Chinnusamy V, Zhu J, Zhu JK: Small RNAs as big players in plant abiotic stress responses and nutrient deprivation. Trends Plant Sci 2007, 12:301-309
- 10. Tang X, Tang G, Ozcan S: Role of microRNAs in diabetes. *Biochim Biophys Acta* 2008, **1779**:697-701.
- Voinnet O: Origin, biogenesis, and activity of plant microRNAs. Cell 2009, 136:669-687.
- Baek D, Villen J, Shin C, Camargo FD, Gygi SP, Bartel DP: The impact of microRNAs on protein output. Nature 2008, 455:64-71.
- Bagga S, Bracht J, Hunter S, Massirer K, Holtz J, Eachus R, Pasquinelli AE: Regulation by let-7 and lin-4 miRNAs results in target mRNA degradation. Cell 2005. 122:553-563.
- Lim LP, Lau NC, Garrett-Engele P, Grimson A, Schelter JM, Castle J, Bartel DP, Linsley PS, Johnson JM: Microarray analysis shows that some microRNAs downregulate large numbers of target mRNAs. Nature 2005, 433:769-773
- Selbach M, Schwanhausser B, Thierfelder N, Fang Z, Khanin R, Rajewsky N: Widespread changes in protein synthesis induced by microRNAs. *Nature* 2008. 455:58-63.
- Yekta S, Shih IH, Bartel DP: MicroRNA-directed cleavage of HOXB8 mRNA. Science 2004, 304:594-596.
- White KP, Rifkin SA, Hurban P, Hogness DS: Microarray analysis of Drosophila development during metamorphosis. Science 1999, 286:2179-2184.
- Goldsmith MR, Shimada T, Abe H: The genetics and genomics of the silkworm, Bombyx mori. Annu Rev Entomol 2005, 50:71-100.
- Aboobaker AA, Tomancak P, Patel N, Rubin GM, Lai EC: Drosophila microRNAs exhibit diverse spatial expression patterns during embryonic development. Proc Natl Acad Sci USA 2005, 102:18017-18022.
- Aravin AA, Lagos-Quintana M, Yalcin A, Zavolan M, Marks D, Snyder B, Gaasterland T, Meyer J, Tuschl T: The small RNA profile during *Drosophila* melanogaster development. Dev Cell 2003, 5:337-350.
- Lagos-Quintana M, Rauhut R, Lendeckel W, Tuschl T: Identification of novel genes coding for small expressed RNAs. Science 2001, 294:853-858.
- Ruby JG, Stark A, Johnston WK, Kellis M, Bartel DP, Lai EC: Evolution, biogenesis, expression, and target predictions of a substantially expanded set of Drosophila microRNAs. Genome Res 2007, 17:1850-1864.
- Stark A, Kheradpour P, Parts L, Brennecke J, Hodges E, Hannon GJ, Kellis M: Systematic discovery and characterization of fly microRNAs using 12 Drosophila genomes. Genome Res 2007, 17:1865-1879.
- 24. Lai EC, Tomancak P, Williams RW, Rubin GM: Computational identification of *Drosophila* microRNA genes. *Genome Biol* 2003, 4:R42.
- Sandmann T, Cohen SM: Identification of novel Drosophila melanogaster microRNAs. PLoS One 2007, 2:e1265.
- Campbell CL, Black WCt, Hess AM, Foy BD: Comparative genomics of small RNA regulatory pathway components in vector mosquitoes. BMC Genomics 2008, 9:425.
- Mead EA, Tu Z: Cloning, characterization, and expression of microRNAs from the Asian malaria mosquito, Anopheles stephensi. BMC Genomics 2008. 9:244.
- 28. Behura SK: Insect microRNAs: Structure, function and evolution. Insect Biochem Mol Biol 2007, 37:3-9.
- Weaver DB, Anzola JM, Evans JD, Reid JG, Reese JT, Childs KL, Zdobnov EM, Samanta MP, Miller J, Elsik CG: Computational and transcriptional evidence for microRNAs in the honey bee genome. Genome Biol 2007, 8: BOT
- 30. Singh J, Nagaraju J: In silico prediction and characterization of microRNAs from red flour beetle (Tribolium castaneum). *Insect Mol Biol* 2008, 17:477, 424
- 31. Wei Y, Chen S, Yang P, Ma Z, Kang L: Characterization and comparative profiling of the small RNA transcriptomes in two phases of locust. *Genome Biol* 2009, 10:R6.
- 32. Cao J, Tong C, Wu X, Lv J, Yang Z, Jin Y: Identification of conserved microRNAs in *Bombyx mori* (silkworm) and regulation of fibroin L chain

- production by microRNAs in heterologous system. *Insect Biochem Mol Biol* 2008. **38**:1066-1071.
- He PA, Nie Z, Chen J, Chen J, Lv Z, Sheng Q, Zhou S, Gao X, Kong L, Wu X, Jin Y, Zhang Y: Identification and characteristics of microRNAs from Bombyx mori. BMC Genomics 2008, 9:248.
- Tong CZ, Jin YF, Zhang YZ: Computational prediction of microRNA genes in silkworm genome. J Zhejiang Univ Sci B 2006, 7:806-816.
- Yu X, Zhou Q, Li SC, Luo Q, Cai Y, Lin WC, Chen H, Yang Y, Hu S, Yu J: The silkworm (Bombyx mori) microRNAs and their expressions in multiple developmental stages. PLoS One 2008. 3:e2997.
- Zhang Y, Zhou X, Ge X, Jiang J, Li M, Jia S, Yang X, Kan Y, Miao X, Zhao G, Li F, Huang Y: Insect-Specific microRNA Involved in the Development of the Silkworm Bombyx mori. PLoS One 2009, 4:e4677.
- Kato M, de Lencastre A, Pincus Z, Slack FJ: Dynamic expression of small non-coding RNAs, including novel microRNAs and piRNAs/21U-RNAs, during Caenorhabditis elegans development. Genome Biol 2009, 10:R54.
- 38. Reddy AM, Zheng Y, Jagadeeswaran G, Macmil SL, Graham WB, Roe BA, Desilva U, Zhang W, Sunkar R: Cloning, characterization and expression analysis of porcine microRNAs. *BMC Genomics* 2009, **10**:65.
- Sunkar R, Zhou X, Zheng Y, Zhang W, Zhu JK: Identification of novel and candidate miRNAs in rice by high throughput sequencing. BMC Plant Biol 2008, 8:25.
- Zhang Y, Zhou X, Ge X, Jiang J, Li M, Jia S, Yang X, Kan Y, Miao X, Zhao G, Li F, Huang Y: A uniform system for microRNA annotation. RNA 2003, 9:277-279
- Glazov EA, Cottee PA, Barris WC, Moore RJ, Dalrymple BP, Tizard ML: A microRNA catalog of the developing chicken embryo identified by a deep sequencing approach. Genome Res 2008, 18:957-964.
- Zhang Y, Zhou X, Ge X, Jiang J, Li M, Jia S, Yang X, Kan Y, Miao X, Zhao G, Li F, Huang Y: A mammalian microRNA expression atlas based on small RNA library sequencing. Cell 2007, 129:1401-1414.
- Morin RD, O'Connor MD, Griffith M, Kuchenbauer F, Delaney A, Prabhu AL, Zhao Y, McDonald H, Zeng T, Hirst M, Eaves CJ, Marra MA: Application of massively parallel sequencing to microRNA profiling and discovery in human embryonic stem cells. Genome Res 2008, 18:610-621.
- Arora A, McKay GJ, Simpson DA: Prediction and verification of miRNA expression in human and rat retinas. *Invest Ophthalmol Vis Sci* 2007, 48:3962-3967.
- Karali M, Peluso I, Marigo V, Banfi S: Identification and characterization of microRNAs expressed in the mouse eye. Invest Ophthalmol Vis Sci 2007, 48:509-515
- Poy MN, Hausser J, Trajkovski M, Braun M, Collins S, Rorsman P, Zavolan M, Stoffel M: miR-375 maintains normal pancreatic alpha- and beta-cell mass. Proc Natl Acad Sci USA 2009, 106:5813-5818.
- Teleman AA, Maitra S, Cohen SM: Drosophila lacking microRNA miR-278 are defective in energy homeostasis. Genes Dev 2006, 20:417-422.
- Lau NC, Lim LP, Weinstein EG, Bartel DP: An abundant class of tiny RNAs with probable regulatory roles in *Caenorhabditis elegans*. Science 2001, 294:858-862.
- Tyler DM, Okamura K, Chung WJ, Hagen JW, Berezikov E, Hannon GJ, Lai EC: Functionally distinct regulatory RNAs generated by bidirectional transcription and processing of microRNA loci. Genes Dev 2008, 22:26-36.
- Bender W: MicroRNAs in the Drosophila bithorax complex. Genes Dev 2008, 22:14-19.
- Stark A, Bushati N, Jan CH, Kheradpour P, Hodges E, Brennecke J, Bartel DP, Cohen SM, Kellis M: A single Hox locus in *Drosophila* produces functional microRNAs from opposite DNA strands. *Genes Dev* 2008, 22:8-13.
- 52. Frasch M: A matter of timing: microRNA-controlled temporal identities in worms and flies. *Genes Dev* 2008, **22**:1572-1576.
- Lu J, Shen Y, Wu Q, Kumar S, He B, Shi S, Carthew RW, Wang SM, Wu Cl: The birth and death of microRNA genes in *Drosophila*. Nat Genet 2008, 40:351-355.
- Animal microRNA target prediction by incorporating diverse sequencespecific determinants.
- 55. Riddiford LM: Juvenile hormone: the status of its "status quo" action. Arch Insect Biochem Physiol 1996, 32:271-286.
- Watanabe K, Hull JJ, Niimi T, Imai K, Matsumoto S, Yaginuma T, Kataoka H: FXPRL-amide peptides induce ecdysteroidogenesis through a G-protein coupled receptor expressed in the prothoracic gland of *Bombyx mori*. Mol Cell Endocrinol 2007, 273:51-58.

- Sheng Z, Ma L, Cao MX, Jiang RJ, Li S: Juvenile hormone acid methyl transferase is a key regulatory enzyme for juvenile hormone synthesis in the Eri silkworm, Samia cynthica ricini. Arch Insect Biochem Physiol 2008, 69:143-154.
- Kamimura M, Takahashi M, Kikuchi K, Reza AM, Kiuchi M: Tissue-specific regulation of juvenile hormone esterase gene expression by 20hydroxyecdysone and juvenile hormone in Bombyx mori. Arch Insect Biochem Physiol 2007. 65:143-151.
- Zechel C: The germ cell nuclear factor (GCNF). Mol Reprod Dev 2005, 72:550-556
- Drevet JR, Swevers L, latrou K: Developmental regulation of a silkworm gene encoding multiple GATA-type transcription factors by alternative splicing. J Mol Biol 1995, 246:43-53.
- Yasukochi Y, Peterson JA, Masters BS: NADPH-cytochrome c (P-450) reductase. Spectrophotometric and stopped flow kinetic studies on the formation of reduced flavoprotein intermediates. J Biol Chem 1979, 254:7097-7104.
- Mach V, Takiya S, Ohno K, Handa H, Imai T, Suzuki Y: Silk gland factor-1 involved in the regulation of *Bombyx* sericin-1 gene contains fork head motif. J Biol Chem 1995, 270:9340-9346.
- Kokubo H, Ueno K, Amanai K, Suzuki Y: Involvement of the Bombyx Scr gene in development of the embryonic silk gland. Dev Biol 1997, 186:46-57.
- Rehwinkel J, Natalin P, Stark A, Brennecke J, Cohen SM, Izaurralde E: Genome-wide analysis of mRNAs regulated by Drosha and Argonaute proteins in Drosophila melanogaster. Mol Cell Biol 2006, 26:2965-2975.
- Nguyen HT, Frasch M: MicroRNAs in muscle differentiation: lessons from Drosophila and beyond. Curr Opin Genet Dev 2006, 16:533-539.
- Sokol NS, Ambros V: Mesodermally expressed *Drosophila* microRNA-1 is regulated by Twist and is required in muscles during larval growth. *Genes Dev* 2005, 19:2343-2354.
- Rao PK, Kumar RM, Farkhondeh M, Baskerville S, Lodish HF: Myogenic factors that regulate expression of muscle-specific microRNAs. Proc Natl Acad Sci USA 2006, 103:8721-8726.
- Zhang Y, Huang J, Jia S, Liu W, Li M, Wang S, Miao X, Xiao H, Huang Y: SAGE tag based cDNA microarray analysis during larval to pupal development and isolation of novel cDNAs in *Bombyx mori*. Genomics 2007. 90:372-379.
- Stark A, Brennecke J, Russell RB, Cohen SM: Identification of Drosophila MicroRNA targets. PLoS Biol 2003, 1:E60.
- Kato M, Paranjape T, Ullrich R, Nallur S, Gillespie E, Keane K, Esquela-Kerscher A, Weidhaas JB, Slack FJ: The mir-34 microRNA is required for the DNA damage response in vivo in C. elegans and in vitro in human breast cancer cells. Oncogene 2009, 28:2419-2424.
- 71. Clevers H: Wnt/beta-catenin signaling in development and disease. *Cell* 2006. **127**:469-480.
- Grigoryan T, Wend P, Klaus A, Birchmeier W: Deciphering the function of canonical Wnt signals in development and disease: conditional loss- and gain-of-function mutations of beta-catenin in mice. Genes Dev 2008, 22:2308-2341.
- Kennell JA, Gerin I, MacDougald OA, Cadigan KM: The microRNA miR-8 is a conserved negative regulator of Wnt signaling. Proc Natl Acad Sci USA 2008, 105:15417-15422.
- Okamura K, Phillips MD, Tyler DM, Duan H, Chou YT, Lai EC: The regulatory activity of microRNA* species has substantial influence on microRNA and 3' UTR evolution. Nat Struct Mol Biol 2008, 15:354-363.
- German MA, Pillay M, Jeong DH, Hetawal A, Luo S, Janardhanan P, Kannan V, Rymarquis LA, Nobuta K, German R, De Paoli E, Lu C, Schroth G, Meyers BC, Green PJ: Global identification of microRNA-target RNA pairs by parallel analysis of RNA ends. Nat Biotechnol 2008, 26:941-946.
- 76. RepBase. http://www.girinst.org.
- 77. Rfam FTP. http://www.sanger.ac.uk/Software/Rfam/ftp.shtml.
- 78. SilkDB. http://silkworm.genomics.org.cn/silkdb/.
- Hofacker IL: Vienna RNA secondary structure server. Nucleic Acids Res 2003, 31:3429-3431.

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