Accepted Manuscript

The *microRNA ame-miR-279a* regulates sucrose responsiveness of forager honey bees (*Apis mellifera*)

Fang Liu, Tengfei Shi, Wei Yin, Xin Su, Lei Qi, Zachary Y. Huang, Shaowu Zhang, Linsheng Yu

PII: S0965-1748(17)30140-6

DOI: 10.1016/j.ibmb.2017.09.008

Reference: IB 2994

To appear in: Insect Biochemistry and Molecular Biology

Received Date: 29 March 2016

Revised Date: 20 August 2017

Accepted Date: 14 September 2017

Please cite this article as: Liu, F., Shi, T., Yin, W., Su, X., Qi, L., Huang, Z.Y., Zhang, S., Yu, L., The *microRNA ame-miR-279a* regulates sucrose responsiveness of forager honey bees (*Apis mellifera*), *Insect Biochemistry and Molecular Biology* (2017), doi: 10.1016/j.ibmb.2017.09.008.

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.





1	
2	The microRNA ame-miR-279a regulates sucrose responsiveness of
3	forager honey bees (Apis mellifera)
4	
5	Fang Liu ^{*1a} , Tengfei Shi ^{1a} , Wei Yin ^b , Xin Su ^a , Lei Qi ^a , Zachary Y. Huang* ^c , Shaowu Zhang ^d ,
6	Linsheng Yu ^a
7	
8	^a College of Animal Science and Technology, Anhui Agricultural University, 230000, Hefei, Anhui,
9	China; ^b The Core Facility, Zhejiang University School of Medicine, Zhejiang University,
10	Hangzhou 310058, China; ^c Department of Entomology, Michigan State University, East Lansing,
11	Michigan, United States of America; ^d Research School of Biology, College of Medicine, Biology
12	and Environment, The Australian National University, Australia.
13	
14	
15	*Correspondence: Zachary Y. Huang, Department of Entomology, Michigan State University, East
16	Lansing, Michigan, United States of America e-mail: bees@msu.edu; Fang Liu, College of Animal
17	Science and Technology, Apiculture Research Institute, Anhui Agricultural University, 230000,
18	Hefei, Anhui, China. Tel.: +86 18297912164; e-mail: lfxiaomifeng@ahau.edu.cn;
19	¹ These authors contributed equally to this work.
20	

21 Abstract

22 Increasing evidence demonstrates that microRNAs (miRNA) play an important role in the 23 regulation of animal behaviours. Honey bees (Apis mellifera) are eusocial insects, with honey bee 24 workers displaying age-dependent behavioural maturation. Many different miRNAs have been 25 implicated in the change of behaviours in honey bees and *ame-miR-279a* was previously shown to 26 be more highly expressed in nurse bee heads than in those of foragers. However, it was not clear 27 whether this difference in expression was associated with age or task performance. Here we show 28 that ame-miR-279a shows significantly higher expression in the brains of nurse bees relative to 29 forager bees regardless of their ages, and that ame-miR-279a is primarily localized in the Kenyon 30 cells of the mushroom body in both foragers and nurses. Overexpression of ame-miR-279a 31 attenuates the sucrose responsiveness of foragers, while its absence enhances their sucrose 32 responsiveness. Lastly, we determined that ame-miR-279a directly target the mRNA of Mblk-1. 33 These findings suggest that *ame-miR-279a* plays important roles in regulating honey bee division 34 of labour. 35 36 Key words: microRNA, Honey bee, proboscis extension reflex, sucrose responsiveness, division

37 of labour.

39 1. Introduction

The honey bee (Apis mellifera. L) is a eusocial insect and a good model organism 40 41 to study the mechanisms and evolution of social behaviours (Robinson et al., 2005). The workers in the colony exhibit age-related division of labour: young honey bees 42 usually engage in within-nest tasks such as brood care ("nursing"), while the old 43 44 honey bees forage outside for different resources (pollen, nectar, water and propolis) (Winston, 1987; Robinson, 1992). However, the division of labour is very flexible: 45 46 bees can accelerate or reverse their behavioural development according to the colony needs (Robinson, 1992; Huang and Robinson, 1996). 47 Numerous studies have focused on the molecular mechanisms underpinning 48 division of labour. Behavioural changes are associated with gene expression changes 49 in the honey bee brain (Whitfield et al., 2003). A number of genes, such as period 50 (Toma et al., 2000), acetylcholinesterase (Shapira et al., 2001), foraging (Ben-shahar 51 et al., 2002, 2005) and malvolio (Ben-shahar et al., 2004) are reported to be involved 52 in the behavioural transition from nurse to forager. MicroRNAs (miRNAs) are 53 54 endogenous small non-coding RNAs (18~24nt) which downregulate gene expression

by mRNA cleavage or translation repression (Bartel, 2004). One single miRNA may

target many mRNAs, and a single mRNA may contain binding sites for many

57 different miRNAs. This leads to a complex regulatory system for biological processes,

such as cell proliferation, differentiation and apoptosis, embryonic development,

neurogenesis, immunity response and disease resistance (Ambros, 2004; Pillai, 2005;

60 Vasudevan, 2007; Legeai et al., 2010).

Several miRNAs were reported to be involved in the honey bee behavioural
maturation process. Behura and Whitfield (2010) found that *miR-276* was upregulated
in young nurses, and had obviously higher expression in young and old nurses than in
young and old foragers, suggesting its involvement in the behavioural maturation

from nurses to foragers. Hori et al. (2010) found that *ame-miR-276* and

66 *ame-miR-1000* are enriched in the optic lobes and in small type Kenyon cells of honey

- bees and that their targets may encode neural function related genes. Greenburg et al.
- (2012) found that *miR-2796* is highly expressed in bee brain, and binds to the coding

region of phospholipase C (PLC)-epsilon gene, which was implicated in neuronal 69 development and differentiation in mammals (Wing et al., 2003), and reported to be 70 transcriptionally regulated in association with division of labour in honey bees 71 (Tsuchimoto et al., 2004). Nunes et al. (2013) identified more than 70 miRNAs that 72 were regulated by the gene vitellogenin, and one of these was ame-miR-279, which 73 may be associated with foraging behavior. Still, the precise mechanism of how 74 miRNAs regulate the division of labour in honey bees is poorly understood. 75 76 Nine miRNAs were previously found to be significantly differentially expressed between nurses and foragers. One of these was ame-miR-279a, which was 77 up-regulated in nurses, and *Mblk-1* was predicted as a candidate target of 78 ame-miR-279a through bioinformatics (Liu et al., 2012). In the present study, we 79 further investigate the role of *ame-miR-279a* in honey bee behavioural development. 80 We show that *ame-miR-279a* is mainly localized in the Kenyon cells of the honey bee 81 mushroom body, and overexpression of ame-miR-279a attenuates the sucrose 82 responsiveness of foragers, while its inhibition enhances their sucrose responsiveness. 83 84 Furthermore, we found that *ame-miR-279a* directly targets the mRNA of *Mblk-1*. 85

86 2. Materials and methods

87 2.1 Honey bees collections

88 European honey bees, Apis mellifera, were maintained according to standard beekeeping practices at Anhui Agriculture University, Hefei, China. Nurses were 89 caught when they had their heads inside cells feeding the larvae. Foragers with 90 91 pollens on their corbiculae were captured at the entrance of the hive. One-day-old 92 honey bees were obtained by removing honeycombs with capped pupae from a typical colony to an incubator (33 °C) until adults emerged. Each one-day-old honey bee was 93 94 marked with a paint dot on the thorax, and kept in the incubator for an hour before being put back into the original colony. A total of 200-300 one-day-old honey bees 95 96 were marked from each typical colony, and three independent typical colonies were used in this study. Three single-cohort colonies were also made, each with about 1000 97 one-day-old honey bees obtained as described, an unrelated mated queen, an empty 98

comb for queen to lay eggs, a comb containing some honey and pollen, all placed in 99 small hive boxes (Whitfield et al., 2003). 100 Twenty 12-day-old nurses (12N) and 30-day-old foragers (30F) were captured 101 respectively from each of the three typical colonies, while another twenty of 102 12-day-old nurses (12N) and 12-day-old ("precocious") foragers (12PF), and 103 30-day-old ("overaged") nurses (30ON) and 30-day-old foragers (30F) were captured 104 from the three single-cohort colonies. The collected honey bees were kept in an 105 106 incubator (33 °C) before their heads were removed for brain dissection to extract RNA for real-time quantitative polymerase chain reaction (RT-qPCR) and northern blot 107

analysis. The honey bees for behavioural experiments were collected from typical

109 colonies. More details are provided later in Section 2.6.

110

111 2.2 Oversupply/inhibition of ame-miR-279a in honey bees

A mimic of *ame-miR-279a* with the sense strand (5'ugacuagauccacacucauuaa3') 112 and the antisense strand 5'aaugaguguggaucuagucauu3') including a 2 nt-3'overhang 113 114 (UU) and 2 nt-5'trim was synthesized by GenPharma (Shanghai, China). An inhibitor (5'uuaaugaguguggaucuaguca3'), a single stranded RNA exactly complementary to 115 ame-miR-279a sequence was also synthesized. A mimic control by using nonsense 116 sequence (sense: 5'uucuccgaacgugucacgutt3', antisense: 5'acgugacacguucggagaatt3') 117 and an inhibitor control using nonsense sequence (5'caguacuuuuguguaguacaa3') were 118 also synthesized. 119

Twenty foragers from a typical colony were used in each treatment and feeding 120 treatments were carried out in three independent experiments. The bees were 121 122 cold-anaesthetized, secured in 0.5-ml Eppendorf tubes with a strip of insulating tape (Supplementary Fig. S1), and kept in an incubator (28 °C, 70% relative humidity) for 123 at least an hour to recover. There were four groups of foragers in the experiment, 124 namely groups fed with the mimic of ame-miR-279a (M), the mimic control of 125 nonsense sequences (NS), the inhibitor of ame-miR-279a (I) and the inhibitor control 126 of nonsense sequences (INS) respectively. Each forager was fed with 10 µl 50% 127 sucrose solution containing 6.6 µg of each synthetic reagent. All the foragers were 128

129	fed to satiety with 50% sucrose solution after treatments (Fig. S2), and kept in the
130	incubator in darkness (28°C, 70% relative humidity). The ame-miR-279a expression
131	in the brains of the foragers was measured 24h after feeding.
132	
133	2.3 RT-PCR and qRT-PCR analysis
134	Bee brains were dissected according to Whitfield et al (2003), then processed for
135	total RNA extraction using a miRNeasy Mini Kit (Qiagen, Germany). The sample
136	quality and quantity were confirmed using a NanoDrop (Thermo Fisher Scientific,
137	Wilmington, DE, USA), and the samples were stored at -80 °C.
138	Total RNA (0.5 μ g per sample) was reverse transcribed with a universal adaptor
139	primer and primeScript RTase. PCR was performed at the same time with specific
140	forward primer (Table 1) and Uni-miR qPCR primer according to the instructions of
141	the SYBR PrimeScript miRNA RT-PCR Kit (TakaRa). The reactions were performed
142	in a TC PCR Thermocycle Instrument (BIOER) under the following conditions: 50 °C
143	for 60 min, 85 °C for 5 s. The qRT-PCR assays were performed in the ABI
144	StepOnePlus TM Real-Time PCR system. Amplification was carried out in 25- μ l
145	reaction volume, containing 10 μ l SYBR premix Ex Taq II, 2 μ l first strand cDNA, 6
146	μ l RNase free water, 0.8 μ l of 10 μ M of each of F and R of the specific primer (Table
147	1). PCR conditions were 95 °C for 30 s, followed by 40 cycles of 95 °C for 5 s and 60
148	°C for 30 s, followed by the melting curve (60 °C -95 °C). β -actin was used as the
149	reference gene. For each gene, test reactions were amplified in quadruplicate along
150	with a no-template and a no-enzyme control. Relative gene expression was calculated
151	using the $2^{-\Delta\Delta^{Ct}}$ method (Livak and Schmittgen, 2001).

152

153 2.4 Northern blot

Total RNA (15 μg per sample) from 20 honey bees brains was separated through a
15% denaturing polyacrylamide gel, then transferred to Hybond-N nylon membranes
by Mini Tans-Blot (Liuyi, Beijing, China) and cross-linked by exposing to ultraviolet
light. DNA oligonucleotides with reverse complementarity to specific sequences were
incorporated with a single digoxigenin-labeled dideoxyuridine-triphosphate

159 (DIG-ddUTP) (Schmitz et al., 1991) by terminal transferase. The sequence of

160 *ame-miR-279a* probe was 5'*uuaaugaguguggaucuaguca3*'. The probe hybridizations

and washes were performed at 65 °C according to the instructions of DIG Northern

162 Starter Kit (Roche, Shanghai, China). Finally, the blots were exposed to Kodak film

according to the method established by Ramkissoon et al. (2006).

164

165 2.5 In Situ Hybridization

166 The honey bee brains were prepared according to Olivier et al. (2008), with the modification that each brain was fixed in 4 % paraformaldehyde (PFA, Sigma) at 4 °C 167 for 30 min, and dehydrated in ascending concentrations of ethanol, embedded in 168 paraffin, then sectioned 10 µm from the frontal side. In situ hybridization was 169 performed according to the kit instructions of BOSTER (# MK10197). The main steps 170 were as follows: the endogenous enzymes in the brain sections were firstly inactivated 171 with 3% H₂O₂; then the sections were treated with pepsin diluted with 3% citric acid 172 for 20 min at room temperature, and washed using PBS; each section was incubated 173 174 with 20 µl hybrid liquid of ame-*miR*-279a probe (5'ttaatgagtgtggatctagtca3') overnight in 40°C; the reactions were blocked and sample incubated with biotinylated 175 anti-mouse digoxin. Colour development was carried out according to the instructions 176 of DAB kit. Finally, sections were dehydrated through a graded series of methanol, 177 soaked with xylene, mounted with neutral gum and examined with a TissueFAXS plus 178 microscope (TissueGnostics, Austria). 179

180

181 2.6 Behavioural experiments

Foragers (N=60-70) were captured from three independent typical colonies, with over 20 foragers per colony. The bees were restrained as mentioned above. The foragers were divided into two groups, one group was fed with *ame-miR-279a* mimic (279a-M), and another one was fed with the mimic control nonsense sequences (279aM-NS). Similarly, another group of foragers (N=60-70) was collected from the same colonies. One half of these foragers were fed with *miR-279a* inhibitor (279aI), another half were fed with the inhibitor control nonsense sequence (279aI-NS). Each

forager was fed with 4.5 µl 50 % sucrose solution containing 1 µg of each synthetic 189 reagent. The foragers were fed to satiety with 50 % sucrose solution after being fed 190 191 the reagents, then put back into the incubator .The bees were tested for sucrose responsiveness using the proboscis extension reflex (PER) assay 24 h and 48 h after 192 treatment. Both antenna of foragers was touched with a droplet of ascending 193 concentrations of sucrose: 0.1, 0.3, 1, 3, 10 and 30 % (w: w) to test their sucrose 194 responsiveness according to previous studies (Pankiw et al., 2001; Page et al., 1998). 195 196 Analysis of variance (ANOVA) was used to analyze the data with PER response as a dependent variable. PER response (%) was analyzed after arcsine-square root 197 transformation. Sugar concentration was treated as a repeated measures variable. 198 Bee brains in the 279aM and 279aM-NS groups were dissected imediately after 199 PER for total RNA extraction according to Section 2.3. The expression of 200 *ame-miR-279a* and *Mblk-1* were quantified using qRT-PCR with β -actin as a control 201 gene (Table 1). 202

203

204 2.7 Western blot

Proteins (90 µg per samples) were extracted from 15 honey bee heads using the 205 Tissue or Cell Total Protein Extraction Kit (Sangon Biotech, Shanghai, China). The 206 protein samples were separated through a 5% denaturing polyacrylamide gel, and 207 208 transferred to nitrocellulose membranes (Pall Life Sciences, Shanghai, China). Non-specific binding-sites on the membranes were blocked with 5 % nonfat milk in 209 TBST for 2 h at room temperature. The membrane was incubated with TBST 210 containing 5 % nonfat milk and diluted rabbit anti-*Mblk-1* polyclonal antibody (1: 200) 211 (SBS, Beijing, China) overnight at 4 °C. It was then washed, incubated with 212 horseradish peroxidase-labeled anti-rabbit IgG (1: 500) (BeyotimeBiotech, Shanghai, 213 China) for an hour at room temperature, and washed again. The immunological 214 detection was carried out according to instructions of the Enhanced HRP-DAB 215

216 Chromogenic Substrate Kit (Tiangen Biotech, Beijing, China).

217

218 2.8 S2 cell culture and luciferase reporter assay

A 421-bp fragment from *Mblk-1* 3'UTR and its mutant sequence and a 249-bp 219 coding region of *ame-miR-279a* were synthesized and amplified using 2×PCR Mix 220 (TaKaRa) (Fig. S3). The Mblk-1 3' UTR and its mutant were cloned into a 221 pAc5.1-firefly luciferase-V5-His vector respectively (Fig. S4A), and the 222 ame-miR-279a coding region was cloned into a pAc5.1-V5-His vector (Fig. S4B), 223 *XhoI* and *NotI* restriction sites were added to the 5' end of the forward and reverse 224 primers, respectively (Table. 2). Drosophila S₂ cells were cultured with 10% fetal 225 226 bovine serum (HyClone) in Schneider's Insect Medium (Invitrogen, Carlsbad, USA). Cells were seeded at 1×10^6 cells per well in a 12-well plate. One day later, 227 ame-miR-279a expression vector (pAc-ame-miR-279a) was co-transfected with either 228 pAc-fluc-Mblk-13'UTR, pAc-fluc-Mblk-13'UTRm, or an empty vector (pAc) in the 229 cells using the calcium phosphate transfection method as described by Tiscornia et al 230 (2006). In all cases, $12 \mu l CaCl_2$ (2 M) and 6 μg transfer vector were mixed, and 1.5 231 µg of pCopia-Renilla luciferase was added as internal control. Forty eight hours after 232 transfection, luciferase assays were performed using a dual-specific luciferase assay 233 234 kit (#RG027, Biyuntian, Shanghai, China). Renilla luciferase activity provided normalization for firefly luciferase activity. 235 236

237 2.9 Statistical analysis

Statistical analysis was conducted as indicated in the text and in figure legends.All t-tests used were two tailed. All tests were done by SPSS 16.0.

240

241 **3. Results**

3.1 The expression of ame-miR-279 paralogs in the brains of nurse and forager bees
We had previously detected a significantly higher expression level of *ame-miR-279a* in the heads of nurses compared to foragers in normal colonies (Liu et al., 2012), and *ame-miR-279b*, *ame-miR-279c*, *ame-miR-279d* were also detected in

- honey bees (Chen et al., 2010; Qin et al., 2014). What might be the differences in
- 240 noney bees (chen et al., 2010, Qin et al., 2017). What highly be the anterences in
- expression among these *miR-279* paralogs between nurses and foragers? As shown in
- Fig.1, there was a significantly higher level of *ame-miR-279a* in the brain of nurses

and foragers than *ame-miR-279b*, *ame-miR-279c*, *ame-miR-279d*. It reveals the
important role of *ame-miR-279a* in the brain function of the honey bee.

251

252 3.2 The expression pattern of ame-miR-279a in the brains of nurses and foragers

There was a significantly higher expression of *ame-miR-279a* in the heads of 253 nurses than in those of foragers in typical colonies (Liu et al., 2012), and it showed a 254 high degree of temporal specificity during the development of adult workers, with the 255 256 highest expression in the 12-day-old nurses and remaining stable in over 30-day-old foragers (Shi et al., 2014). These suggest a possible important function of 257 ame-miR-279a in honey bee behavior plasticity. To confirm this hypothesis, the 258 expression and localization of ame-miR-279a in the brains of nurses and foragers 259 were investigated. We first measured the *ame-miR-279a* expression in the brains of 260 nurses and foragers exhibiting normal behavior in typical colonies. A t-test showed 261 that ame-miR-279a was significantly highly expression in 12-day-old nurses 262 compared to the 30-day-old foragers (t=3.79, P<0.05) (Fig. 2A). However, the 263 264 differential expression of *ame-miR-279a* between nurses and foragers may be associated with their ages but not their different behavior. To resolve this question, we 265 created the single-cohort colonies, and tested ame-miR-279a expression in foragers 266 and nurses of the same age. As expected, this pattern stayed the same regardless 267 268 whether nurses and foragers were both young (12 days old) or both old (30 days old) in single cohort colonies (Fig. 2B). The ame-miR-279a expression between nurses of 269 different ages (12 vs. 30 days old) was not significantly different, nor was it between 270 foragers of different ages (Fig. S5). Northern blot further confirmed that 271 272 ame-miR-279a had a higher expression in nurses than in foragers, regardless of whether both groups were 12 days old or 30 days old (Fig. 2C). 273 To determine the localization of ame-miR-279a in adult honey bee brains (nurses 274

and foragers), in situ hybridization was performed using LNA (locked nucleic acid)
miRNA. The results showed that *ame-miR-279a* (brown staining) was predominantly
expressed in the Kenyon cells of the mushroom bodies (Fig. 3A, B) and in the lamina

of the optic lobes in nurse and forager (Fig. 3A, C). The blank control produced no

brown staining (Fig. 3D). Moreover, ame-miR-279a expression in the brain showed 279 no obvious spatial difference between the nurse bees and forager bees even when they 280 were of the same age (Fig. S6). Taken all together, these results confirmed the 281 important role of *ame-miR-279a* in the bee behavioral maturation. 282 283 3.3 inhibition and overexpression of ame-miR-279a in the honey bee 284 Considering the importance of *ame-miR-279a* in behavioral maturation, we 285 286 decided to overexpress and inhibit the miRNA in honey bees to examine possible effects on behavior. The synthetic inhibitor (anti-miRNA) and mimic of 287 ame-miR-279a were fed to foragers together with 50% sucrose solution. The 288 qRT-PCR confirmed the overexpression and inhibition of *ame-miR-279a* in the brains 289 290 of honey bee in the presence of the mimic and inhibitor respectively. As shown in Fig. 4, the *ame-miR-279a* expression in foragers from the M group was significantly 291 higher than in the NS group, while ame-miR-279a expression in foragers from the I 292 group was significantly lower than that of the INS group. 293 294 3.4 ame-miR-279a affects the sucrose responsiveness of foragers 295 To further investigate the possible function of ame-miR-279a in the honey bees' 296

behavioral maturation, we tested the effect of *ame-miR-279a* on PER first by using a 297 298 mimic. As was no significant difference in PER between 24 and 48 h (F=3.22, df = 1, 48; P = 0.08), we analyzed the two sets of data together. PER response varied 299 significantly with sugar concentrations (F=15.78, df = 5, 48; P<0.001). PER response 300 was significantly lower in bees fed with a mimic (279aM) compared to a control 301 group fed with nonsense control (279aM-NS) (F=13.12, df=1, 5; P < 0.001, Fig. 5A). 302 We then tested the effect of *ame-miR-279a* on PER by using its inhibitor. There 303 was no significant difference in PER between 24 and 48 h (F=1.07, df = 1, 48; P > 304 0.1), and we analyzed the two sets of data together. PER response varied significantly 305 with sugar concentrations (F=14.71, df = 5, 48; P<0.001). PER response was 306 307 significantly higher in bees fed with an inhibitor (279aI) compared to a control group fed with nonsense control (279aI-NS) (F=4.96, df=1, 5; P < 0.04, Fig. 5B). 308

309	
310	3.5 Quantification of the expression of ame-miR-279a and Mblk-1
311	Mblk-1 was predicted as the target of ame-miR-279a (Liu et al., 2012). In order
312	to confirm their interaction, we detected the expression of <i>ame-miR-279a</i> and <i>Mblk-1</i>
313	in the brains of honey bees from the experimental foragers above. As expected,
314	ame-miR-279a had much higher expression in the brains of foragers in group 279aM
315	than in group 279aM-NS (t=14.924, P<0.05) (Fig. 6), while Mblk-1 had significantly
316	lower expression in the brains of foragers from the 279aM group than from the
317	279aM-NS group (t=3.884, P<0.05) 24 h after treatment (Fig. 6). The <i>Mblk-1</i> protein
318	level in forager heads from the corresponding honey bees was further examined by
319	western blot, as shown in Fig. 6. Honey bees in 279aM group showed a lower Mblk-1
320	protein level than the 279aM-NS group 24h after treatment (Fig. 6). Similar results
321	were obtained 48 h after treatment (Fig. S7).
322	
323	3.6 Confirmation of the interaction of ame-miR-279a with Mblk-1 using a luciferase
324	reporter assay
324 325	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned
324 325 326	<i>reporter assay</i> To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted
324 325 326 327	<i>reporter assay</i> To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated
324 325 326 327 328	<i>reporter assay</i> To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> 'UTR (Fig. 8A). A sequence with mutations (m) was also
324 325 326 327 328 329	<i>reporter assay</i> To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3' UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> ' UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as
324 325 326 327 328 329 330	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-1</i> 3'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-1</i> 3'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a
324 325 326 327 328 329 330 331	<i>reporter assay</i> To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> 'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-13</i> 'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i>
324 325 326 327 328 329 330 331 332	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> 'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-13</i> 'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i> was co-transfected with pAc-fluc- <i>Mblk-13</i> 'UTR in S ₂ cells, the luciferase activity
324 325 326 327 328 329 330 331 332 333	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> 'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-13</i> 'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i> was co-transfected with pAc-fluc- <i>Mblk-13</i> 'UTR in S ₂ cells, the luciferase activity significantly decreased compared to the assay involving co-transfection with
324 325 326 327 328 329 330 331 332 333 334	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-1</i> 3'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-1</i> 3'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i> was co-transfected with pAc-fluc- <i>Mblk-1</i> 3'UTR in S ₂ cells, the luciferase activity significantly decreased compared to the assay involving co-transfection with pAc-fluc- <i>Mblk-1</i> 3'UTR m and pAc (t=10.07, P<0.0001, Fig. 8B). Moreover,
324 325 326 327 328 329 330 331 332 333 334 335	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3' UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> ' UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-13</i> ' UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i> was co-transfected with pAc-fluc- <i>Mblk-13</i> ' UTR in S ₂ cells, the luciferase activity significantly decreased compared to the assay involving co-transfection with pAc-fluc- <i>Mblk-13</i> ' UTR m and pAc (t=10.07, P<0.0001, Fig. 8B). Moreover, <i>ame-miR-279a</i> expression directly reduced the <i>Mblk-1</i> mRNA and protein levels (Fig.
324 325 326 327 328 329 330 331 332 333 334 335 336	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> 'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-13</i> 'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i> was co-transfected with pAc-fluc- <i>Mblk-13</i> 'UTR in S ₂ cells, the luciferase activity significantly decreased compared to the assay involving co-transfection with pAc-fluc- <i>Mblk-13</i> 'UTR m and pAc (t=10.07, P<0.0001, Fig. 8B). Moreover, <i>ame-miR-279a</i> expression directly reduced the <i>Mblk-1</i> mRNA and protein levels (Fig. 4). All these results support the conclusion that <i>Mblk-1</i> is a direct target of
324 325 326 327 328 329 330 331 332 333 334 335 336 337	reporter assay To test whether <i>ame-miR-279a</i> actually targets the <i>Mblk-1</i> 3' UTR, we subcloned a 421-bp fragment of the 3'UTR region of <i>Mblk-1</i> mRNA that included the predicted <i>ame-miR-279a</i> recognition site (Fig. 7) into a luciferase reporter plasmid designated as pAc-fluc- <i>Mblk-13</i> 'UTR (Fig. 8A). A sequence with mutations (m) was also constructed as the negative control for the same reporter assay, named as pAc-fluc- <i>Mblk-13</i> 'UTR-m. The coding region of <i>ame-miR-279a</i> was cloned into a pAc5.1-V5-His vector designated as pAc- <i>ame-miR-279a</i> . When pAc- <i>ame-miR-279a</i> was co-transfected with pAc-fluc- <i>Mblk-13</i> 'UTR in S ₂ cells, the luciferase activity significantly decreased compared to the assay involving co-transfection with pAc-fluc- <i>Mblk-13</i> 'UTR m and pAc (t=10.07, P<0.0001, Fig. 8B). Moreover, <i>ame-miR-279a</i> expression directly reduced the <i>Mblk-1</i> mRNA and protein levels (Fig. 4). All these results support the conclusion that <i>Mblk-1</i> is a direct target of <i>ame-miR-279a</i> .

339 4. Discussion

The role of miRNA in insect behavior has been well established in recent years 340 341 (Lucas and Raikhel, 2013). The miR-iab4/iab8 locus controls self-righting behavior in larvae of *Drosophila* by repressing the Hox gene *Ultrabithorax* (Picao-Osorio et al., 342 2015). Ecdysone controls let-7 - Complex to repress the circadian gene clockwork 343 orange to regulate the circadian rhythms of Drosophila (Chen et al., 2014). 344 MicroRNA-133 inhibits the behavioral aggregation of locusts by controlling dopamine 345 346 (Yang et al., 2014). *MicroRNA-932* regulates the memory of honey bee by targeting actin (Alexandre et al., 2014). Dme-miR-279 regulates the JAK/STAT pathway to 347 drive the rest: activity rhythms in *Drosophila* (Luo and Sehgal, 2012). In this study, 348 we concentrated on *ame-miR-279a* since its expression was significantly higher in 349 nurses than that of foragers, and showed a high degree of temporal specificity in 350 typical colonies (Liu et al., 2012; Shi et al., 2014). However, it was not clear whether 351 the expression of ame-miR-279a was associated with task performance (nursing) or 352 age (young bees). We decoupled the task performance and age in honey bees by using 353 354 single cohort colonies, a method regularly used to accomplish this (e.g. Robinson et al. 1989, Ben-Shahar et al., 2002). We determined that the ame-miR-279a expression was 355 always higher in nurses than in foragers regardless of whether they were young 356 (typical nurses vs. precocious foragers), or were both old (overaged nurses vs. typical 357 foragers). These results are consistent with another study in honey bees, in which the 358 foraging gene was shown to regulate the behavioral transition between nurses and 359 foragers (Ben-Shahar et al., 2002). Thus, we deduced that there is a good correlation 360 between ame-miR-279a and honey bee behavioral changes. 361

Mushroom bodies (MBs) are higher-order brain centres thought to be important for sensory integration, learning and memory formation in the honey bee (Giurfa, 2007; Menzel, 1999; Menzel, 2012). MBs have a high degree of structural plasticity depending on caste and task performance, suggesting that they are associated with honey bee social behaviours (Robinson et al., 1997; Withers et al., 1993). The MBs are famous as important brain regions of olfactory learning in the vinegar fly, *Drosophila melanogaster* (Hayashi et al., 2009). It has been reported that

dme-miR-279 was detected with strongest expression in the head epidermis in regions 369 adjacent to where the sensory organ progenitors form in Drosophila (Stark et al., 370 2005). A putative orphan receptor (HR38) homologue that mediates 371 ecdysteroid-signaling, showed higher expression in the MBs of forager brains 372 compared to nurse bees, suggesting its involvement in regulation of the division of 373 labour of the workers (Yamazaki et al., 2006). In this study, we demonstrated that 374 ame-miR-279a is expressed more in the Kenyon cells of the mushroom bodies, 375 376 suggesting that *ame-miR-279a* may play a role in social behaviour. However, there were no obvious spatial differences between nurses and foragers when we used in situ 377 hybridization. This suggests that the differences in ame-miR-279a levels between 378 nurses and foragers detected with RT-qPCR may represent increased expression in the 379 same cells. This is consistent with the expression pattern of the *foraging* gene in nurse 380 and forager bees, which was proved to regulate the division of labour of honey bees 381 (Ben-Shahar et al., 2002). 382 It was reported that *dme-miR-279* can regulate the formation of carbon dioxide 383 384 (CO₂) neurons by targeting the transcription factor Nerfin-1 in Drosophila (Cayirlioglu et al., 2008), and that *Prospero* restricts CO₂ neuron formation indirectly 385 via miR-279 and directly by repressing the common targets, Nerfin-1 and Esg, 386 suggesting the importance of *dme-miR-279* in the neuron and olfactory system 387 388 development in Drosophila (Hartl et al., 2011). In this study, we found that overexpression of *ame-miR-279a* attenuated the sucrose responsiveness of foragers 389 (Fig. 5A), while its reduction enhanced their sucrose responsiveness (Fig. 5B). 390

Responsiveness to sucrose is associated with foraging choices, as bees with high

392 sucrose responsiveness preferentially collect pollen or water while bees with low

sucrose responsiveness mainly collect nectar (Pankiw and Page, 1999; Scheiner et al.,

2001a), suggesting the importance of *ame-miR-279a* in regulating honey bee olfactory

behaviour. Moreover, we found that nurses always had higher expression of

ame-miR-279a than foragers regardless of their age (Fig. 2). It has been demonstrated

that nurse bees are less responsive than foragers to gustatory stimuli (Scheiner et al.,

2001a,b), and water foragers have higher responsiveness to sucrose than both of

399 pollen and nectar foragers (Pankiw, 2005). In our study, overexpression of ame-miR-279a in foragers may make them physiologically similar to nurses, resulting 400 in lower sucrose responsiveness (Fig. 5A, B), and suggesting that *ame-miR-279a* may 401 modulate the honey bee behavioural transition from nurses to foragers, or stimulate 402 foragers to change their behaviour from nectar collection to water or pollen foraging 403 when colony conditions demand so. 404 We have previously predicted *Mblk-1* to be a possible target for *ame-miR-279a* 405 406 (Liu et al., 2012). The expression of ame-miR-279a is largely confined to the mushroom body of the honey bee brain (Fig. 3), and overexpression of *ame-miR-279a* 407 significantly inhibited the mRNA and protein expression of *Mblk-1* in forager brains 408 (Fig. 6). Moreover, our luciferase assay confirmed that ame-miR-279a targets the 409 3'UTR of Mblk-1 because transfection of pAc-fluc-Mblk-13'UTR reduced the 410 luciferase activity and pAc-fluc-Mblk-13'UTRm rescued this suppression to the same 411 level as that of the blank control (Fig. 8). These results strongly indicate that 412 ame-miR-279a directly targets Mblk-1. The Mblk-1 gene, encoding a putative 413 414 transcription factor is also expressed preferentially in the large-type Kenyon cells of honey bee MBs. It contains several motifs characteristic of transcription factors, 415 including RHF1 and RHF2, a nuclear localization signal and glutamine-run motifs 416 (Takeuchi et al., 2001). Thus, Mblk-1 is thought to be involved in brain function by 417 regulating transcription of its target genes. It has been reported that Mblk-1 may 418 function in MB neural circuits directly modulated by the Ras/MAPK pathway (Park et 419 al., 2003). E93, a homologue of Mblk-1 in Drosophila, expressed highly in the brain 420 of the fly, has been shown to affect olfactory sensory neurons (Jafari et al., 2012). 421 422 MBR-1, another homologue of Mblk-1 in the nematode *Caenorhabditis elegans*, was also reported to have neuronal functions, in which it is required for the pruning of 423 specific neurites that occur during larval development (Kage et al., 2005). Moreover, 424 it was also shown that MBR-1 is required for olfactory plasticity in adult animals 425 (Hayashi et al., 2009; Takayanagi-Kiya et al., 2017). Taken together, we deduce that 426 Mblk-1 may be involved in the regulation of behavioral plasticity of honey bee 427 through its target gene ame-miR-279a in the MBs. 428

- 429 In summary, we found that *ame-miR-279a* showed significantly higher expression
- 430 in nurses than in foragers regardless of their ages, and *ame-miR-279a* was primarily
- 431 localized in the Kenyon cells of the mushroom body of foragers and nurses;
- 432 overexpression of *ame-miR-279a* attenuated the sucrose responsiveness of foragers,
- 433 while its inhibition enhanced their sucrose responsiveness. Moreover, we determined
- 434 that *ame-miR-279a* directly targets the mRNA of *Mblk-1*. These findings suggest that
- 435 *ame-miR-279a* plays important roles in regulating honey bee division of labour.
- 436

437 Acknowledgements

This work was supported by grants of National Natural Science Foundation of
China (31302039), Education Department Research Project of Anhui Province
(2013SQRL018ZD). We thank Wenfeng Chen for his kindly provide with luciferase
reporter plasmid, and thank Tiande Liang for technical assistance in collecting honey
bees and preparing samples, Aung Si, Zhiguo Li and Shoujun Huang for critically
reading the manuscript.

444

445 Author's contributions

- 446 F.L. planned the experiments, performed *In Situ Hybridization*, the reporter assay,
- data analysis and wrote the manuscript. T.F.S. performed RNA extraction, RT-PCR
- and qRT-PCR *analysis*, western blot. W.Y., X.S. and L.Q. performed experimental
- 449 experiments. Z.Y.H. was involved in experimental design, data analysis and
- 450 manuscript revision. S.W.Z. and L.S.Y. performed manuscript revision. All authors
- 451 have read the final draft of the manuscript

453 **References**

- 454 Ambros, V., 2004. The functions of animal microRNAs. Nature 431, 350–355.
- Bartel, D. P., 2004. MicroRNAs: genomics, biogenesis, mechanism, and function. Cell 116,
 281–297.
- 457 Behura, S.K. and Whitfield, C.W., 2010. Correlated expression patterns of microRNA genes with

458 age-dependent behavioural changes in honeybee. Insect Mol. Biol. 19, 431–439.

- Ben-Shahar, Y., 2005. The foraging gene, behavioral plasticity, and honeybee division of labor. J.
 Comp. Physiol. A 191: 987–994
- 461 Ben-Shahar, Y., Dudek, N.L. and Robinson, G.E., 2004. Phenotypic deconstruction reveals

462 involvement of manganese transporter malvolio in honey bee division of labor. J. Exp. Biol.
463 207, 3281–3288.

- Ben-Shahar, Y., Robichon, A., Sokolowski, M.B. and Robinson, G.E., 2002. Influence of gene
 action across different time scales on behavior. Science 296, 741–744.
- Cayirlioglu, P., Kadow, I. G., Zhan, X., Okamura, K., Suh, G. S. B., Gunning, D., Lai, E. C., and
 Zipursky, S. L., 2008. Hybrid Neurons in a MicroRNA Mutant Are Putative Evolutionary

468 Intermediates in Insect CO2 Sensory Systems. Science 319, 1256–1260.

469 Chen, X., Yu, X., Cai, Y., Zheng, H., Yu, D., Liu, G., Zhou, Q., Hu, S., and Hu, F. 2010.

- 470 Next-generation small RNA sequencing for microRNAs profiling in the honey bee Apis
 471 mellifera. Insect Mol. Biol. 19, 799-805.
- 472 Cristino, A. S., Barchuk, A. R., Freitas, F.C.P., Narayanan, R. K., Biergans, S. D., Zhao, Z. et al.,

473 2014. Neuroligin-associated microRNA-932 targets actin and regulates memory in the
474 honeybee. Nat. Commun. 5, 5529.

475 Giurfa, M., 2007. Behavioral and neural analysis of associative learning in the honeybee: a taste

476 from the magic well. J Comp Physiol A Neuroethol. Sens Neural. Behav. Physiol. 193,
477 801–824.

- Greenberg, J.K., Xia, J., Zhou, X., Thatcher, S.R., Gu, X., Ament, S.A. et al., 2012. Behavioral
 plasticity in honey bees is associated with differences in brain microRNA transcriptome.
 Genes Brain Behav, 11, 660–670.
- 481 Hartl, M., Loschek, L. F., Stephan, D., Siju, K. P., Knappmeyer, C., and Kadow, I. C. G., 2011. A
- 482 New Prospero and microRNA-279 Pathway Restricts CO2 Receptor Neuron Formation. J.

- 483 Neurosci. 31, 15660–15673.
- Hayashi, Y., Hirotsu, T., Iwata, R., Kage-Nakadai, E., Kunitomo, H., Ishihara, T., Iino, Y., and
 Kubo, T. 2009. A trophic role for Wnt-Ror kinase signaling during developmental pruning in *Caenorhabditis elegans*. Nat. Neurosci. 12, 981-987.
- 487 Heisenberg, M. Mushroom body memoir: From maps to models.Nat. Rev. Neurosci.2003, 4,
 488 266–275.
- 489 Hori, S., Kaneko, K., Saito, T. H., Takeuchi, H., and Kubo, T., 2010. Expression of two
- 490 microRNAs, ame-mir-276 and -1000, in the adult honeybee (*Apis mellifera*) brain.
 491 Apidologie.
- 492 Huang, Z.Y. and Robinson, G.E., 1996. Regulation of honey bee division of labor by colony age
- 493 demography. Behav. Ecol. Sociobiol. 39, 147–158.Kage, E., Hayashi, Y., Takeuchi, H.,
- Hirotsu, T., Kunitomo, H., Inoue, T., Arai, H., Iino, Y., and Kubo, T. (2005). MBR-1, a novel
 helix-turn-helix transcription factor, is required for pruning excessive neurites in *Caenorhabditis elegans. Current Biology* 15,1554-1559.
- Legeai, F., Rizk, G., Walsh, T., Edwards, O., Gordon, K., Lavenier, D., et al., 2010. Bioinformatic
 prediction, deep sequencing of microRNAs and expression analysis during phenotypic
 plasticity in the pea aphid, Acyrthosiphon Pisum. BMC Genomics 11, 281–290.
- 500 Liu, F., Peng, W., Li, Z.G., Li, W.F., Li, L., Pan, J. et al., 2012. Next-generation small RNA
- sequencing for microRNAs profiling in *Apis mellifera*: comparison between nurses and
 foragers. Insect Mol. Biol. 21, 297–303.
- 503 Livak, K.J. and Schmittgen, T.D., 2001. Analysis of relative gene expression data using real-time 504 quantitative PCR and the $2^{-\Delta\Delta Ct}$ method. Methods 25, 402–408.
- 505 Lucas, K., and Raikhel, A. S. (2013). Insect MicroRNAs: Biogenesis, expression profiling and
- 506 biological functions. Insect Mol. Biol. 43, 24–38.
- 507 Luo, W. and Sehgal, A., 2012. Regulation of Circadian Behavioral Output via a
- 508 MicroRNA-JAK/STAT Circuit. Cell 148, 765–779.
- 509 Menzel, R., 1999. Memory dynamics in the honeybee. J. Comp. Physiol. A 185, 323–340.
- 510 Menzel, R., 2012. The honeybee as a model for understanding the basis of cognition.Nat. Rev.
- 511 Neurosci. 13, 758–768.

- 512 Nunes, F.M.F., Ihle, K.E., Mutti, N.S., Simões, Z.L.P. and Amdam, G.V., 2013. The gene
- 513 vitellogenin affects microRNA regulation in honey bee (*Apis mellifera*) fat body and brain. J.

514 Exo. Biol. 216, 3724–3732.

- Olivier, V., Massou, I., Celle, O., Blanchard, P., Schurr, F., Ribière, M., Gauthier, M., 2008. In
 situ hybridization assays for localization of the chronic bee paralysis virus in the honey bee
 (*Apis mellifera*) brain. J. Virol. Methods 153, 232–237.
- 518 Page, R. J., Erber, J. and Fondrk, M., 1998. The effect of genotype on response thresholds to
- sucrose and foraging behavior of honey bees (*Apis mellifera* L.). J. Comp. Physiol. A 182,
 489–500.
- 521 Pankiw, T., 2005. The honey bee foraging behavior syndrome: quantifying the response threshold
- 522 model of division of labor. IEEE Xplore Conference: Swarm Intelligence Symposium, DOI:
 523 10.1109/SIS.2005.1501595
- 524 Pankiw, T. and Page Jr, R.E., 1999. The effect of genotype, age, sex, and caste on response
- thresholds to sucrose and foraging behavior of honey bees (*Apis mellifera* L.). J. Comp.
- 526 Physiol. A 185, 207–213.
- 527 Pankiw, T., Waddington, K.D. and Page, R.E., 2001. Modulation of sucrose response thresholds in
- honey bees (*Apis mellifera* L.): influence of genotype, feeding, and foraging
 experience. J. Comp. Physiol. A 187, 293–301.
- 530 Park, J. M., Kunieda, T., and Kubo, T., 2003. The activity of Mblk-1, a nushroom body-selective
- transcription factor from the honeybee, is modulated by the Ras/MAPK pathway. J. Biol.
- 532 CHEM. 278, 18689–18694.
- 533 Picao-Osorio, J., Johnston, J., Landgraf, M., Berni, J., and Alonso, C. R. (2015).
- 534 MicroRNA-encoded behavior in Drosophila. *Science* 350, 815–820.
- Pillai, R. S., 2005. MicroRNA function: Multiple mechanisms for a tiny RNA? *Rna* 11,
 1753–1761.
- Qin, Q.-H., Wang, Z.-L., Tian, L.-Q., Gan, H.-Y., Zhang, S.-W., and Zeng, Z.-J. (2014). The
 integrative analysis of microRNA and mRNA expression inApis melliferafollowing
 maze-based visual pattern learning. Insect Sci. 21, 619-636.
- 540 Ramkissoon, S.H., Mainwaring, L.A., Sloand, E.M., Young, N.S. and Kajigaya, S., 2006.
- 541 Nonisotopic detection of microRNA using digoxigenin labeled RNA probes. Mol. Cell.

- 542 Probe 20, 1–4.
- Robinson, G.E., Fahrbach, S.E. and Winston, M.L., 1997. Insect societies and the molecular
 biology of social behavior. Bioessays 19, 1099–1108.
- Robinson, G.E., Grozinger, C.M. and Whitfield, C.W., 2005. Sociogenomics: social life in
 molecular terms. Nat. Rev. Genet. 6, 257–270.
- 547 Robinson, G.E., 1992. Regulation of division of labor in insect societies. Annu. Rev. Entomol. 37,
 548 637–665.
- Robinson, G. E., and Page, R. E., 1989. Genetic determination of nectar foraging, pollen foraging,
 and nest-site scouting in honey bee colonies. Behav. Ecol. Sociobiol. 24, 317–323.
- 551 Scheiner, R., Page, R.E. and Erber, J., 2001a. The effects of genotype, foraging role and sucrose
- perception on the tactile learning performance of honey bees (*Apis mellifera* L.). Neurobiol.
- 553 Learn. Mem. 76, 138–150.
- 554 Scheiner, R., Page, R.E. and Erber, J., 2001b. Responsiveness to sucrose affects tactile and
- olfactory learning in preforaging honey bees of two genetic strains. Behav. Brain Res. 120,
 67–73.
- 557 Schmitz, G. G., Walter, T., Seibl, R., and Kessler, C., 1991. Nonradioactive labeling of
- 558oligonucleotides in vitro with the hapten digoxigenin by tailing with terminal transferase.
- 559 Analytical Biochemistry 192, 222-231.
- Shapira, M., Thompson, C.K. Soreq, H. and Robinson, G.E., 2001. Changes in neuronal
 acetylcholinesterase gene expression and division of labor in honey bee colonies. J. Mol.
 Neurosci. 17, 1–12.
- Shi, T., Liu, F., Yu, L., Wang, T. and Qi, L., 2014. Expression levels of three miRNAs in the brain
 of different day-old workers of *Apis mellifera ligustica* (Hymenoptera: Apidae). Acta
- 565 Entomological Sinica 57, 1368–1374.
- 566 Stark, A., Brennecke, J., Bushati, N., Russell, R. B., and Cohen, S. M., 2005. Animal MicroRNAs
- 567 Confer Robustness to Gene Expression and Have a Significant Impact on 3'UTR Evolution.
 568 Cell 123, 1133–1146.
- 569 Takayanagi-Kiya, S., Kiya, T., Kunieda, T., and Kubo, T. 2017. Mblk-1 Transcription Factor
- 570 Family: Its Roles in Various Animals and Regulation by NOL4 Splice Variants in Mammals.
- 571 Int. J. Mol. Sci. 18, 246.

- 572 Takeuchi, H., Kage, E., Sawata, M., Kamikouchi, A., Ohashi, K., Ohara, M. et al., 2001.
- 573 Identification of a novel gene, Mblk-1, that encodes a putative transcription factor expressed
- preferentially in the large-type Kenyon cells of the honeybee brain, Insect Mol. Biol. 10,
 487–494.
- 576 Tiscornia, G., Singer, O., and Verma, I. M., 2006. Production and purification of lentiviral vectors.
 577 Nat. Protoc. 1, 241–245.
- Toma, D.P., Moore, D., Bloch, G. and Robinson, G.E., 2000. Changes in period expression in the
 brain and division of labor in honey bee colonies. PNAS 97, 6914–6919.
- 580 Tsuchimoto, M., Aoki, M., Takada, M., Kanou, Y., Sasagawa, H., Kitagawa, Y. & Kadowaki, T.,
- 581 2004. The changes of gene expression in honeybee (*Apis mellifera*) brains associated with
 582 ages. Zoolog. Sci. 21, 23–28.
- 583 Vasudevan, S., Tong, Y., and Steitz, J. A., 2007. Switching from Repression to Activation:
- 584 MicroRNAs Can Up-Regulate Translation. Science 318, 1931–1934.
- 585 Whitfield, C.W., Cziko, A.M. and Robinson, G.E., 2003. Gene expression profiles in the brain
 586 predict behavior in individual honey bees. Science 302, 296–299.
- 587 Winston, M.L., 1987. The biology of the honeybee. Havard University Press, Cambridge, MA.
- 588 Withers, G.S., Fahrbach, S.E. and Robinson, G.E., 1993. Selective neuroanatomical plasticity and
 589 division of labour in the honeybee. Nature 364, 238–240.
- 590 Wing, M.R., Bourdon, D.M. & Harden, T.K., 2003. PLC-epsilon: a shared effector protein in Ras-,
- 591 Rho-, and G alpha beta gamma-mediated signaling. Mol. Interv. 3, 273–280.
- Yamazaki. Y., Shirai, K., Paul, R.K., Fujiyuki, T., Wakamoto, A., Takeuchi, H. and Kubo, T., 2006.
 Differential expression of HR38 in the mushroom bodies of the honeybee brain depends on
 the caste and division of labor. Febs. Lett. 580, 2667–2670.
- Yang, M., Wei, Y., Jiang, F., Wang, Y., Guo, X., He, J., and Kang, L. (2014). MicroRNA-133
 Inhibits Behavioral Aggregation by Controlling Dopamine Synthesis in Locusts. PLOS Genet.
 10, e1004206.
- 598

```
Table 1 Primer sequences used for qRT-PCR validation of ame-miR-279a and Mblk-1.
```

- 600
- **Table 2** Primer sequences used for RT-PCR amplification of 3'UTR and

pri-miR-279a.

Table 1

Primer	5' to 3'
Mblk-1 -F	AACACCAAATACGACCCAAAAC
Mblk-1 -R	CAACAGAGCCTTCTCCACTTCT
ame-miR-279a-F	CTTTCTAAGTATCAATAATG
ame-miR-279aR	TCTTAAAATTCATATTCATA
β-actin-F β-actin-R	TGCCAACACTGTCCTTTCTG AGAATTGACCCACCAATCCA
Table 2	
Primer	5' to 3'
Mblk-1 3'UTR-F	CGCCCGAAACCGCGAAAGAA
Mblk-1 3'UTR-R	GACGTCGAATCACGCCTTGT
pri-miR-279a-F	CTTTCTAAGTATCAATAATG
pri-miR-279aR	TCTTAAAATTCATATTCATA

608 Figure captions:

- Fig. 1. Expression levels of four *miR-279* paralogs (miR-279a, miR-279b, miR-279c,
 miR-279d) in the brains of nurses and foragers.
- **Fig. 2.** Expression levels (+SE) of *ame-miR-279a* in the brain of 12 and 30 days old
- age-matched nurses and foragers from regular colonies (A) and single-cohort colonies
- (B). Student t-test results were shown, with * denoting P<0.05 and ** denoting
- 614 P<0.01. Data based on three replicates (colonies). (C) Northern blot analysis of
- 615 *ame-miR-279a* in brains of age-matched 12-day-old young nurses (12N) and young
- 616 ("precocious") foragers (12PF), and age-matched 30 days old foragers (30F) and old
- 617 ("overage") nurses (30ON) from single-cohort colonies. 5s rRNA was used as a

618 reference.

619

Fig. 3. Expression of *ame-miR-279a* in the honey bee brain. OL, optic lobe; KC,

621 Kenyon cells. *ame-miR-279a* is highly expressed in the Kenyon cells of the

mushroom bodies and in the lamina of the optic lobes (brown colour) with the

623 positive probe (A). No brown labeling was seen in sections probed with a blank

control (D). Squares delineate regions in shown magnified in BC and EF. There were
no obvious spatial differences between nurses and foragers; these images are from a

626 627 nurse brain.

Fig. 4. *Ame-miR-279a* expression in the brains of foragers after oral feeding with *mimic-mir-279a* (M) or nonsense sequence (NS), or *inhibitot-mir-279a* (I) or inhibitor nonsense sequence (INS). An independent t-test result is shown, data represent the mean from three independent experiments \pm s.e.m * means P<0.05, ** means P<0.01.

632

Fig. 5. Mean score (% +SE) of bees responding to proboscis extension response to

various sugar concentrations after bees treated with a mimic (A) or inhibitor (B) of

635 *ame-miR-279a*. The effect of *ame-miR-279a* on foragers' responsiveness to sucrose.

- Responsiveness to sucrose was significantly lower (P < 0.01) in foragers fed on a
- 637 *miR-279a mimic* (279aM) compared to those fed with a nonsense sequence

638	(279aM-NS). Conversely, response to sucrose was significantly ($P < 0.01$) enhanced
639	in foragers fed on a miR-279a inhibitor (279aI) compared to those fed with a
640	nonsense sequence (279aI-NS). Data from three colonies were analyzed after
641	arsine-square root transformation but presented here without transformation.
642	
643	Fig. 6 . Relative expression levels (\pm SE) of <i>ame-miR-279a</i> and <i>Mblk-1</i> from group
644	279aM and 279aM-NS at 24h after treatment. Student t-test results are shown with *
645	denoting P<0.05, ** denoting P<0.01. Data are from three replicates (colonies).
646	Western blot analysis of Mblk-1 protein in foragers' heads from 279aM and
647	279aM-NS at 24h after treatment, β -actin was used as the reference protein.
648	
649	Fig. 7. Sequences of the interaction sites between <i>ame-miR-279a</i> and <i>Mblk-1-3'UTR</i> .
650	Asterisks indicate mutated site, mutated nucleotide bases are shown in bold. Grey
651	shaded areas indicate canonical 7mer "seed" region that aligns with the target site, the
652	vertical lines indicate contiguous Watson-Crick pairing.
652 653	vertical lines indicate contiguous Watson-Crick pairing.
652 653 654	vertical lines indicate contiguous Watson-Crick pairing.Fig. 8. (A) A schematic representation of the principle behind the luciferase assay.
652 653 654 655	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i>'UTR resulted in dramatic suppression of the
652 653 654 655 656	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i>'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with
652 653 654 655 656 657	vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i> 'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with \pm s.e.m.
652 653 654 655 656 657 658	vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i> 'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with \pm s.e.m.
652 653 654 655 656 657 658 659	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i>'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with ±s.e.m. Fig. S1. The 0.5 ml Eppendorf tube used in our experiment, A: A normal tube; B: The
652 653 654 655 656 657 658 659 660	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i>'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with ±s.e.m. Fig. S1. The 0.5 ml Eppendorf tube used in our experiment, A: A normal tube; B: The tube without lid; C: The tube with the part of top margin removed. C was used in the
652 653 655 656 657 658 659 660 661	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i> 'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with ±s.e.m. Fig. S1. The 0.5 ml Eppendorf tube used in our experiment, A: A normal tube; B: The tube without lid; C: The tube with the part of top margin removed. C was used in the present study. D: Prepared tubes inside a rack with a pair of forceps and insulation. E:
652 653 654 655 656 657 658 659 660 661 662	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i> 'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with ±s.e.m. Fig. S1. The 0.5 ml Eppendorf tube used in our experiment, A: A normal tube; B: The tube without lid; C: The tube with the part of top margin removed. C was used in the present study. D: Prepared tubes inside a rack with a pair of forceps and insulation. E: A restrained forager.
 652 653 655 656 659 660 661 662 663 	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i> 'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with ±s.e.m. Fig. S1. The 0.5 ml Eppendorf tube used in our experiment, A: A normal tube; B: The tube without lid; C: The tube with the part of top margin removed. C was used in the present study. D: Prepared tubes inside a rack with a pair of forceps and insulation. E: A restrained forager.
 652 653 654 655 657 658 659 660 661 662 663 664 	 vertical lines indicate contiguous Watson-Crick pairing. Fig. 8. (A) A schematic representation of the principle behind the luciferase assay. (B) co-transfection of <i>pAc-fluc-Mblk-13</i> 'UTR resulted in dramatic suppression of the luciferase activity. A normalized firefly/renilla luciferase value was plotted with ±s.e.m. Fig. S1. The 0.5 ml Eppendorf tube used in our experiment, A: A normal tube; B: The tube without lid; C: The tube with the part of top margin removed. C was used in the present study. D: Prepared tubes inside a rack with a pair of forceps and insulation. E: A restrained forager. Fig. S2. During the hour of recovery, the honey bees were fed with 10 µl sucrose

total darkness. After the test, the honey bees were fed 50 μ l sucrose solution (50%, in

 dH_2O and put back into the incubator to wait for the next test.

668	
669	Fig. S3. A 421-bp fragment from <i>Mblk-1</i> 3'UTR (A) and a 249-bp coding region of
670	<i>ame-miR-279a</i> (B).
671	
672	Fig. S4. Schematic overview of the pAc5.1-firefly luciferase-V5-His vector used to
673	construct pAc-fluc-Mblk-13'UTR plasmids (A), and the pAc5.1-V5-His vector used
674	to construct pAc-ame-miR-279a plasmid (B).
675	
676	Fig. S5. Mean (+SE) levels of <i>ame-miR-279a</i> expression in nurses and foragers. There
677	were no statistical differences between different ages. Data based on three replicates.
678	(Data are mean \pm s.e.m, n=3).
679	
680	Fig. S6. Expression of <i>ame-miR-279a</i> in normal forager brain (A), 12 days old nurse
681	brain (D), 12 days old forager brain (H). OL, optic lobe; KC, Kenyon cells. Squares
682	delineate regions in shown magnified in BC, EF and IJ. Ame-miR-279a is highly
683	expressed in the Kenyon cells of the mushroom bodies and in the lamina of the optic
684	lobes (brown colour) with the positive probe.
685	
686	Fig. S7. Relative expression levels (+SE) of <i>ame-miR-279a</i> and <i>Mblk-1</i> from group
687	279aM and 279aM-NS at 48h after treatment. Student t-test results are shown with *
688	denoting P<0.05, ** denoting P<0.01. Data are from three replicates (colonies).
689	Western blot analysis of Mblk-1 protein in foragers' heads from 279aM and
690	279aM-NS at 48h after treatment, β -actin was used as the reference protein.
691	
692	Y



Relative expression levels

Ψ















Α



ω

- 1. *Ame-miR-279a* shows significantly higher expression in the brains of nurse bees relative to forager bees regardless of their ages.
- 2. *Ame-miR-279a* is primarily localized in the Kenyon cells of the mushroom body in both foragers and nurses.
- 3. Overexpression of *ame-miR-279a* down-regulates the sucrose responsiveness of foragers, while its absence up-regulates their sucrose responsiveness.
- 4. Ame-miR-279a directly targets the mRNA of Mblk-1.

CERTIFIC AND