Genetic variation in the effect of monoamines on female mating receptivity and oviposition in the adzuki bean beetle, *Callosobruchus chinensis* (Coleoptera: Bruchidae)

Takashi Yamane

**Abstract**

**Background:** Female mate choice after mating is a strong force in sexual selection and could lead to coevolution of mating traits between the sexes. How females of different genotypes respond to substances in the male ejaculate should be mediated by females' mate choices. Monoamines regulate animal physiology and behavior, including the post-mating behavior of females of the adzuki bean beetle, *Callosobruchus chinensis* (Coleoptera: Bruchidae). This study examined differences in females' response to four monoamines (dopamine, octopamine, tyramine, serotonin) between strains from different populations of *C. chinensis*.

**Results:** Injection with either octopamine or tyramine, two kinds of monoamines significantly reduced female receptivity in two strains with low remating frequencies. None of the four monoamines reduced female receptivity in one strain with high remating frequencies. However, all monoamines reduced it in another strain with high remating frequencies. Oviposition was activated by tyramine on days 1–5 or by serotonin on days 4 and 5 in the two strains with low remating frequencies, but only on day 1 or day 4 in the strains with high remating frequencies.

**Conclusion:** These differences in female response to monoamines, especially tyramine and serotonin, correspond with results of previous studies. They indicate differences in female response to male substances that reduce receptivity and activate oviposition. These findings suggest relationships between the differences in female response to male substances among populations and mutations in the pathways of monoamine biosynthesis or transmission, which in turn determine female mate choice in response to male substances.

**Keywords:** Female mate choice, Seminal fluids, Mating receptivity, Oviposition, Biogenic amines, Adzuki bean beetle

**Background**

The lifetime number of matings and fecundity of females differ among animal species [1-3]. Males can improve their reproductive success by preventing multiple matings by females, increasing the number of offspring produced, and hastening female oviposition, as a result of reduced sperm competition with other males [4]. However, females can improve their reproductive success by mating with multiple males, leading to increased offspring viability [5], despite the costs of mating [6,7]. Furthermore, although male ejaculate has a cost [8], eggs are much more valuable to females than sperm are to males [9], so the quality of mates and the effect on offspring quality would be more important for females than for males; thus, females are thought to select males or their sperm so they can adjust the number of offspring sired by a particular males or overall [2,10].

On this basis, the optimal number of matings and offspring per mating for females might differ from those for males, resulting in sexual conflict over mating traits [11], and cause antagonistic adaptations and counter-adaptations between male and female traits [12,13]. However, male reproductive traits that impose costs for female fitness can be selected by female mate choice, leading to
coevolution between both sexes, if the benefits to females exceed the costs. Because of the importance of similarity and diversity for the genetics of a species, much research has been devoted to antagonistic adaptations or counter-adaptations via sexual conflict and to female mate choice and coevolution of the sexes in terms of traits related to the mating rate and the number of offspring [10,14-19].

Mating can change female behaviors, such as activating oviposition and reducing mating receptivity [4,10]. Such changes are often caused by substances in the male seminal fluids [20-22] that are shown to increase the reproductive success of the males by manipulating females [4]. However, female fecundity and remating rates can differ among populations of a species, possibly reflecting genotypic differences that could be explained by sexual selection via female mate choice based on genotypic differences in the quality of male ejaculate [23-26]. However, no studies have defined the physiological components of seminal substances that reflect these variations and cause differences among female genotypes that would lead to sexual selection via female mate choice.

Monoamines are neuroactive substances that affect behavioral and physiological traits in both vertebrates and invertebrates, and they act as neurotransmitters and endocrine disruptors in the central and peripheral nervous systems [27-29]. They have several known roles in determining the mating status and behavior of invertebrates: in Drosophila melanogaster, octopamine and tyramine regulate egg laying [30-32]; in Caenorhabditis elegans, serotonin (5-HT) regulates egg laying [33]; in female Locusta migratoria, octopamine [34] and tyramine [35,36] modulate muscle contraction in the spermatheca and oviduct, and serotonin [37] modulates it in the spermatheca, suggesting roles in egg laying; and in Lepidoptera, octopamine and analogs increase egg laying [38], serotonin or its metabolites induce mate-rejection behavior [39], and tyramine reduces the production of sex pheromones [40] within females.

In the adzuki bean beetle, Callosobruchus chinensis, male-derived substances reduce female receptivity to mating and activate female oviposition, but the responses of females to these substances and the effects of males differ among strains from different geographic populations [41,42]. The effects of four artificially injected monoamines (dopamine, octopamine, tyramine, and serotonin) on female receptivity and oviposition in one strain of C. chinensis (jC-S) suggested a relationship between these monoamines and female post-mating behavior [43]. Here, I built on this research by comparing the effects of monoamines on female mating receptivity and oviposition among four C. chinensis strains derived from populations from different geographic locations, adding three new strains of C. chinensis (isC, akC02, and mC; Additional file 1: Table S1) to examine any differences between strains in response to monoamines.

**Results**

**Female receptivity**

Generalized linear model (GLM) results showed significant effects of monoamine treatment on female receptivity and on the interaction between female strain and monoamine treatment, but not of female strain (Table 1). A separate GLM analysis revealed a significant effect of monoamine treatment on female receptivity for each strain except isC (jC-S, likelihood ratio $\chi^2 = 39.510, P < 0.0001$; isC, $\chi^2 = 8.826, P = 0.0656$; akC02, $\chi^2 = 38.923, P < 0.0001$; and mC, $\chi^2 = 16.178, P = 0.0028$). In jC-S, octopamine and tyramine significantly reduced female receptivity compared with the control, dopamine, and serotonin (Figure 1). In isC, no differences were significant. In akC02, all monoamines significantly reduced receptivity compared with the control, and tyramine reduced it compared with dopamine, octopamine, and serotonin. In mC, octopamine and tyramine reduced receptivity compared with the control, and tyramine reduced it compared with dopamine. In addition, there was difference between strain in receptivity of control ($\chi^2 = 8.565, P = 0.0357$), but not any between two strains.

**Number of eggs laid by females**

Repeated-measures two-way analysis of variance (ANOVA) revealed that all between-subject sources of variance (monoamine treatment, female strain, treatment × strain) and all within-subject sources of variance (days after injection, days × treatment, days × strain, days × treatment × strain) significantly affected the number of eggs laid (Table 2). Because of the significant interaction effects, separate one-way ANOVA tests were applied to the differences among monoamines in each strain and on each day (Additional file 1: Table S2). Relative to the control, tyramine significantly ($P < 0.05$) increased the number of eggs from day 1 and serotonin from day 4 in jC-S (Figure 2A) and mC (Figure 2D); octopamine did so from day 1 and tyramine on day 4 in isC (Figure 2B); and tyramine did so on day 1 in akC02 (Figure 2C). Results of monoamines on female receptivity and number of eggs laid by females are

Table 1 Results of a generalized linear model testing the effects of strain and monoamine treatment on female receptivity

| Deviance | Source | df | Likelihood ratio $\chi^2$ | P |
|----------|--------|----|--------------------------|---|
| 705.645  | –      | 570 | –                        | – |
| –        | Treatment | 4  | 86.270                  | < 0.0001 |
| –        | Strain     | 3  | 7.711                   | 0.0524  |
| –        | Strain × treatment | 12 | 21.295                 | 0.0462  |
summarized in Table 3. Law dates of them were attached (Additional file 2).

**Discussion**

Artificial injection to *C. chinensis* strain jC-S females revealed a relationship between biogenic monoamines and female mating receptivity, oviposition, or both [43]. My results confirm that result and reveal differences among *C. chinensis* populations in the effects of monoamines on female receptivity and the number of eggs laid. In other studies, oral application of serotonin induced mate-rejection behavior in female *Pieris rapae* and increased the level of serotonin and its metabolites in the brains of females [38]; injection of tyramine reduced production of the sex pheromone bombykol in mated *Bombyx mori* females [39]; and octopamine and analogs applied to the dorsal surface of female *Plodia interpunctella* increased the number of eggs laid [40]. However, this is the first report to show differences among or within populations, which could reflect differences among genotypes, in the effect of monoamines on females of different strains of the same species.

Octopamine and tyramine significantly reduced female receptivity in jC-S and mC compared with the control (Figure 1). Females of both strains remated less frequently than those of the other strains (Additional file 1: Table S1). A previous study [41] showed that two kinds of male-derived substances reduced female receptivity: low-molecular-weight (MW) substances (<3 kDa), which act quickly (within 3–5 h and 1 day after injection), and higher-MW substances (>14 kDa), which act more slowly (2–4 days after injection). The jC-S and mC females responded more strongly than the other females to male substances that reduce female receptivity: jC-S females responded to the low-MW substances within 3 to 5 h and 1 day, and to the higher-MW substances by 2 and 4 days after injection, and mC females responded to the low-MW substances by 1 day and to the higher-MW substances at 2 and 4 days after injection [41]. In contrast, isC females, with a higher remating frequency (Additional file 1: Table S1), responded more weakly to male substances than the other strains: they did not respond to the low-MW substances, and responded to the higher-MW substances at 2 days after injection [41]. The receptivity of isC females was not reduced by the injection of any tested monoamine (Figure 1). Therefore, the effects of biogenic monoamines in these strains coincided with the degree of response to male substances. Likewise, akC02 females, which also respond more weakly than the other strains to

**Table 2 Results of repeated-measures two-way ANOVA of eggs laid by females of four strains in the monoamine injection experiment**

| Source               | df | F     | P     |
|----------------------|----|-------|-------|
| Between-subject      |    |       |       |
| Treatment            | 4  | 8.746 | < 0.0001 |
| Strain               | 3  | 15.593| < 0.0002 |
| Treatment × strain   | 12 | 1.948 | 0.0274 |
| Error                | 19 | 5.389 |       |
| Within-subject       |    |       |       |
| Day                  | 1.92 | 4888.741 | < 0.0001* |
| Day × Treatment      | 7.67 | 5.503  | < 0.0001* |
| Day × strain         | 5.77 | 13.821 | < 0.0001* |
| Day × treatment ×    | 23.06 | 2.140  | 0.0014* |
| Error                | 36.51 | 4.543  |       |

*Corrected P, Greenhouse-Geisser ε = 0.4804.
male substances, did not respond to low-MW substances, and responded to high-MW substances at 2 and 4 days after injection [41]; however, they responded strongly to all four monoamines by reducing their receptivity (Figure 1). Female receptivity can also be reduced by other factors, such as mechanical stimulation during mating, insertion of the male’s aedeagus, mechanical pressure on stretch receptors in the female bursa, and the presence of sperm in the female spermatheca [21]. These factors may be more important than the composition of the seminal fluids in the reduced receptivity of akC02.

Tyramine significantly increased the number of eggs laid on days 1 to 5 and serotonin on days 4 and 5 by jC-S and mC (Figure 2A, D). Tyramine significantly increased the number laid on day 4 by isC females (Figure 2B) and on day 1 by akC02 females (Figure 2C). Octopamine significantly increased the number laid on days 1 to 5 by isC females (Figure 2B). Male-derived substances (>14 kDa) activate female oviposition, and females of jC-S and mC had stronger responses to them than females of isC and

![Figure 2](image_url)

**Figure 2** Effects of monoamines on female oviposition. Total number of eggs laid by *C. chinensis* females of strains (A) jC-S, (B) isC, (C) akC02, and (D) mC after injection with dopamine, octopamine, tyramine, serotonin, or Milli-Q water (control) over 5 days after injection. The number of eggs laid differed significantly among monoamine treatments on days 1 to 5 in jC-S, isC, and mC, and on days 1 and 5 (but not on days 2 to 4) in akC02 (Additional file 1: Table S2). Bars labeled with an asterisk differ significantly from the control within a strain (*P* < 0.05, Dunnett’s test). The numbers in parentheses show the sample size.

| Trait | Strain | Dopamine | Octopamine | Tyramine | Serotonin |
|-------|--------|----------|------------|----------|-----------|
| Receptivity | jC-S | ● | ● | ● | ● |
| | isC | ● | ● | ● | ● |
| | akC02 | ● | ● | ● | ● |
| | mC | ● | ● | ● | ● |
| Egg laying | jC-S | 1-5 d | 4-5 d | ● | ● |
| | isC | 1-5 d | 4 d | ● | ● |
| | akC02 | 1-5 d | 4-5 d | ● | ● |
| | mC | 1-5 d | 4-5 d | ● | ● |

*●* indicates an effect; number indicate days on which effects on egg laying were found.

---

Table 3 Summary of effects of monoamines on reduction of female mating receptivity and eggs laid compared with control of each strain of *Callosobruchus chinensis*

---

Yamane BMC Evolutionary Biology 2014, 14:172
http://www.biomedcentral.com/1471-2148/14/172
akC02; jC-S and mC females responded to these substances derived from males of each of the same four strains used here, but isC females did not respond to substances from three of the strains and akC02 females did not respond to any [42]. Thus, differences among the strains in the numbers of eggs laid coincided with the responses to male substances. The similarities and differences in the effects of the four monoamines, especially tyramine and serotonin, might be related to the effects of male substances.

Serotonin receptors have been identified in invertebrates [44]. Behavioral pharmacology and genetic studies of insect serotonin receptors revealed the specific involvement of receptor subtypes in modulating behaviors [45-48]. Furthermore, because C. elegans mutants of serotonin receptor genes SER-1 and SER-2 stopped laying in response to exogenous serotonin, and expression of SER-1 in vulval muscles recovered the response to serotonin, serotonin acts on egg laying via the SER-1 receptor-coupled signaling pathway in vulval contraction [33]. Thus, the coincidences between the difference of the effects of serotonin on oviposition and the seminal fluid substances that stimulate oviposition among populations indicate that serotonin receptors are involved in responses to male substances that induce female oviposition in C. chinensis. Moreover, tyramine acts as an intermediate precursor for octopamine biosynthesis, but it also has diverse direct roles in physiology and behavior [28,29]. It is possible that octopamine converted from the injected tyramine affected female mating behavior suggested by previous study [43]. Therefore, similarities and differences in male substances related to increased oviposition, and differences among strains, may be related to similarities and differences among the genes involved in the biosynthesis pathway from tyrosine via tyramine to octopamine, which is mediated by the enzymes tyrosine decarboxylase and tyramine-β-hydroxylase [30-32]. They may also be related to differences in the receptors for octopamine, tyramine, or both [28,29], as isC females increased oviposition significantly after injection with octopamine (Figure 2B).

Females of the red flour beetle, Tribolium castaneum, that mated with males of the same genotype remated earlier [26] and their lifetime fecundity was lower [25] than females that mated with males of different genotypes, and females of Musca domestica that mated with males of the same genotype oviposited less than females that mated with males of different genotypes [23]. These studies suggest that females resist male manipulations that reduce their receptivity and induce their fecundity following mating, and that antagonistic coevolution occurs between the sexes as a result of female choice. In contrast, Callosobruchus maculatus females that mated with males of different genotypes remated earlier and their fecundity was lower than females that mated with males of the same genotype [24]. This indicates that coevolution between the sexes resulted from female resistance to male manipulations that reduce their receptivity and induce fecundity.

In addition to female responses, there were differences in the effects of these substances between male genotypes: jC-S male substances more strongly reduced female receptivity and activated oviposition [41,42]. As mentioned above, jC-S females had stronger responses to male substances. Therefore, these results suggest that the ability of males to reduce receptivity and induce fecundity is selected for by the female, and that coevolution may have occurred between the sexes. These male manipulations appear to disadvantage the females—for example, male-derived substances reduce female longevity [42]—but might select for males with a strong ability to secure their paternity by enhancing their male offspring’s fitness, or might select for abilities linked to other traits that enhance benefits to females, such as nutrient availability or offspring viability [14,17].

Conclusion

The four biogenic monoamines produced different responses in the four female strains of C. chinensis. The differences in responses (reduced receptivity or increased oviposition) suggest the result of mutations in the biosynthesis pathways or the receptors for these monoamines. That is, choice among traits is mediated via such mutations. By examining variations between populations in the monoamine composition of male ejaculate, it should be possible to reveal details of the physiological mechanisms that underlie the differences among the female genotypes in their responses to differences in male ejaculate. These differences may lead to sexual selection as a result of female choice and the evolution of the substances in male seminal fluids as a result of coevolution between the sexes.

Ethics

This study had been conducted under the ARRIVE guidelines, which are intended to improve the reporting of animal research on vertebrates or any regulated invertebrates.

Methods

Insects

The four strains of C. chinensis were maintained in a growth chamber at 25°C, 60% relative humidity, and a light–dark cycle of 16 L:8 D. Strains jC-S and mC were reared at the University of Tokyo [49,50] and strains isC and akC02 at Okayama University [51,52]. Approximately 100 adults of each strain were kept in separate plastic Petri dishes (height, 1.5 cm; diameter, 9.1 cm) at Okayama University. In April 2011, the beetles were transferred to a chamber with the same conditions at the National Agricultural Research Center, Tsukuba, Japan. All beetles were reared from eggs laid by randomly collected females on
adzuki bean (Vigna angularis ‘Dainagon’) seeds (restricted to one egg per seed). Each seed was then transferred into a separate well of a 48-well tissue culture plate (Greiner Bio-One, Frickenhausen, Germany) and kept within the chamber described above. Beetles eclosed within the bean and then emerged from it. Thus, virgin male and female adults were collected from the beans in the wells. Emerging females have mature eggs in their oviducts and bursa copulatrix [43].

Injection of monoamines
Dopamine, octopamine (Nacalai Tesque, Kyoto, Japan), tyramine, or serotonin (Sigma-Aldrich, Tokyo, Japan) was dissolved in Milli-Q water to a final concentration of 10% (dopamine and octopamine, 0.53 M; tyramine, 0.58 M) or 2% (serotonin, 0.05 M). Serotonin does not dissolve well in water, so it was not possible to create a concentration greater than 2%.

Virgin females aged 1 to 4 days old were chilled on ice for a few minutes and fixed to agarose medium at room temperature by using fine forceps. Under a microscope, a hole was made between the second and fifth segments of the ventral abdomen with the forceps, and 0.05 μl of a monoamine solution or Milli-Q water (control) was injected through an ultra-fine glass capillary connected to an oil-pressure injector (Nanoject Auto-Nanoliter Injector, Drummond Scientific Company, Broomall, PA, USA).

At 3 to 5 h after the injection, the female and a virgin male of strain jC-S were placed in a small plastic Petri dish (height, 1.5 cm; diameter, 3.0 cm) in the chamber described above. The pair was then observed for 1 h to see whether the female mated.

To count the number of eggs laid, I immediately transferred the mated females into separate wells of a 24-well tissue culture plate (Nalge Nunc International K.K., Tokyo, Japan) and supplied one adzuki bean per female. I counted the eggs oviposited on each bean every day for 5 days. Data for females that died during that period were excluded from analysis. If eggs were laid on a bean, the bean was replaced with a new one.

Cumulative total numbers of eggs (transformed as log \( \log(n + 1) \)) were compared using repeated-measures two-way ANOVA with “treatment” and “strain” as the between-subject factors and “days after injection” as the within-subject factor. Because Mauchly’s test of sphericity indicated a significant violation of the assumption of sphericity (\( P < 0.0001 \)), significance levels for within-subject effects were calculated using a Greenhouse-Geisser correction of the degrees of freedom [54]. When a significant interaction effect was encountered among between-subject or within-subject sources of variance (Table 2), separate one-way ANOVAs were applied to the differences among strains. To compare variables between control and monoamines, when significant effects were detected in the separate one-way ANOVA, Dunnett’s test was used for multiple comparisons at the 5% significance level. All statistical analyses were performed in JMP v. 11.0.0 software [55].

**Additional files**

**Additional file 1: Table S1.** Rearing history and female remating frequency of each strain of Callosobruchus chinensis. Table S2. Results of one-way ANOVAs testing the effects of monoamine treatment on the number of eggs laid by females in each strain at 1 to 5 days after injection.

**Additional file 2: Date of mating receptivity.** Date of number of eggs laid by females.

**Abbreviations**

5-HT: Serotonin; GLM: Generalized linear model; ANOVA: Analysis of variance; MW: Molecular-weight.

**Competing interests**

The author has no financial or non-financial competing interests.

**Acknowledgements**

I thank Dr. Göran Arnqvist, members of his laboratory (Uppsala University) for helpful discussion in the preparation of the manuscript, and two anonymous reviewers for their valuable comments to the manuscript. I also thank Drs. Takahisa Miyatake (Okayama University) and Masako Katsuki (Tsukuba University) for providing the C. chinensis strains. This work was partly supported by Research Fellowships for Young Scientists (Japan Society for the Promotion of Science 233967) to T.Y.

Received: 17 April 2014 Accepted: 23 July 2014

**References**

1. Ridley M. Mating frequency and fecundity in insects. Biol Rev 1988, 63:509–549.
2. Birkhead TR. Promiscuity: An Evolutionary History of Sperm Competition and Sexual Conflict. London: Faber & Faber; 2000.
3. Simmons LW. The evolution of polyandry: sperm competition, sperm selection, and offspring viability. Annu Rev Ecol Evol Syst 2005, 36:125–146.
4. Simmons LW. Sperm Competition and its Evolutionary Consequences in the Insects. Princeton: Princeton University Press; 2001.
5. Slatyer RA, Mautz BS, Patricia R, Backwell Y, Jennions MD. Estimating genetic benefits of polyandry from experimental studies: a meta-analysis. Biol Rev 2012, 87:1–33.
6. Chapman T, Liddle LF, Kalb JM, Wolfner MF, Partridge L. Cost of mating in Drosophila melanogaster females is mediated by male accessory gland products. Nature 1995, 373:241–244.
7. Eady PE, Hamilton L, Lyons RE: Copulation, genital damage and early death in Callosobruchus maculatus. Proc R Soc Lond B Biol Sci 2007, 274:247–252.
8. Dewsbury DA: Ejaculate cost and male choice. Am Nat 1982, 119:501–610.
9. Bateman AJ: Intra-sexual selection in Drosophila. Heredity 1948, 2:349–368.
10. Eberhard WG: Female Control: Sexual Selection by Cryptic Female Choice. Princeton, NJ: Princeton University Press; 1996.
11. Arnqvist G, Rowe L: Sexual Conflict. Princeton, NJ: Princeton University Press; 2005.
12. Holland B, Rice WR: Chase-away sexual selection: antagonistic seduction versus resistance. Evolution 1998, 52:1–7.
13. Lessells CM: The evolutionary outcome of sexual conflict. Philos Trans R Soc Lond B Biol Sci 2006, 361:301–317.
14. Eberhard WG, Cordero C: Sexual selection by cryptic female choice on male seminal products - a new bridge between sexual selection and reproductive physiology. Trends Ecol Evol 1995, 10:493–495.
15. Gavrelles S, Arnqvist G, Friberg U: The evolution of female mate choice by sexual conflict. Proc R Soc Lond B Biol Sci 2001, 268:531–539.
16. Cordero C, Eberhard WG: Female choice of sexually antagonistic male adaptations: a critical review of some current research. J Evol Biol 2003, 16:1–46.
17. Cordero C, Eberhard WG: Interaction between sexually antagonistic selection and mate choice in the evolution of female responses to male traits. Evol Ecol 2005, 19:111–122.
18. Kokko H, Brooks R, Jennions MD, Marley J: The evolution of mate choice and mating biases. Proc R Soc Lond B Biol Sci 2003, 270:653–664.
19. Kvarnemo C, Simmons LW: Polyandry as a mediator of sexual selection before and after mating. Philos Trans R Soc Lond B Biol Sci 2013, 368:20120042.
20. Gillett C: Male accessory gland secritions: modulators of female reproductive physiology and behavior. Ann Rev Entomol 2003, 48:163–184.
21. Wedell N: Female receptivity in butterflies and moths. J Exp Biol 2005, 208:3433–3440.
22. Aula FW, Sirot UK, LaFlamme BA, Rubinstein CD, Wolfner MF: Insect seminal fluid proteins: identification and function. Ann Rev Entomol 2011, 56:21–40.
23. Andrès JA, Arnqvist G: Genetic divergence of the seminal signal-receptor system in houseflies: the footprints of sexually antagonistic coevolution? Proc R Soc Lond B Biol Sci 2001, 268:399–405.
24. Brown DV, Eady PE: Functional incompatibility between the fertilization systems of two allopatric populations of Callosobruchus maculatus (Coleoptera: Bruchidae). Evolution 2001, 55:2257–2262.
25. Nilsson T, Fricke C, Arnqvist G: Patterns of divergence in the effects of mating on female reproductive performance in flour beetles. Evolution 2002, 56:111–120.
26. Nilsson T, Fricke C, Arnqvist G: The effects of male and female genotype on variance in male fertilization success in the red flour beetle Tribolium castaneum. Behav Ecol Sociobiol 2003, 53:227–233.
27. Liebert F, Pilzger HJ: Monoamines and the orchestration of behavior. Bioscience 2004, 54:17–25.
28. Roeder T: Tyramine and octopamine: ruling behavior and metabolism. Ann Rev Entomol 2005, 50:447–477.
29. Lange AB: Tyramine: From octopamine precursor to neuroactive chemical in insects. Gen Comp Endocrinol 2009, 162:18–25.
30. Monastirioti M, Linn CE, White K: Characterization of Drosophila Tyramine β-hydroxylase gene and isolation of mutant flies lacking octopamine. J Neurosci 1996, 16:3900–3911.
31. Monastirioti M: Distinct octopamine cell population residing in the CNS abdominal ganglion controls ovulation in Drosophila melanogaster. Dev Biol 2003, 264:38–49.
32. Cole SH, Carney GI, McClung CA, Willard SS, Taylor BJ, Hirsh J: Two functional but non-complementing Drosophila tyrosine decarboxylase genes: distinct roles for neural tyramine and octopamine in female fertility. J Biol Chem 2005, 280:14948–14955.
33. Denspemy CM, Mackeren SM, Gargou A, Blanco G, Sae JF: Serotonin (5HT), fluoxetine, imipramine and dopamine target distinct 5HT receptor signaling to modulate Caenorhabditis elegans egg-laying behavior. Genetics 2005, 169:1425–1436.
34. Clark J, Lange AB: Octopamine modulates spermathecal muscle contractions in Locusta migratoria. J Comp Physiol A 2003, 189:105–114.
35. Donini A, Lange AB: Evidence for a possible neurotransmitter/neuromodulator role of tyramine on the locust oviducts. J Insect Physiol 2004, 50:351–361.
36. da Silva R, Lange AB: Tyramine as a possible neurotransmitter/neuromodulator at the spermatheca of the African migratory locust, Locusta migratoria. J Insect Physiol 2006, 54:1306–1313.
37. Clark J, Lange AB: The association of serotonin with the spermatheca of the locust, Locusta migratoria. Biochim Biophys Acta 2007, 1777:43–62.
38. Obara Y, Fukano Y, Watanabe K, Ozawa G, Sasaki K: Serotonin-induced mate rejection in the female cabbage butterfly, Pieris rapae crucivora. Naturwisenschaften 2011, 98:989–993.
39. Hiroshina A, Yamaji H, Yoshizawa T, Kusano E, Eto M: Effect of tyramine and stress on sex-pheromone production in the pre- and post-mating silkworm moth, Bombyx mori. J Insect Physiol 2007, 53:1242–1249.
40. Rajasekhar A: Enhanced oviposition in the moth, Plodia interpunctella, after treatment with anajogenic analogues. J Stored Prod Res 2004, 40:331–339.
41. Yamane T, Miyatake T: Evolutionary correlation between male substances and female remating frequency in a seed beetle. Behav Ecol 2012, 23:723–728.
42. Yamane T: Intra-specific variation in the effect of male seminal substances on female oviposition and longevity in Callosobruchus chinensis. Evol Ecol Res 2013, 15:133–140.
43. Yamane T, Miyatake T: Reduced female mating receptivity and activation of oviposition in two Callosobruchus species due to injection of biogenic amines. J Insect Physiol 2010, 56:271–276.
44. Watanabe T, Sadamoto H, Aonuma H: Identification and expression analysis of the genes involved in serotonin biosynthesis and transduction in the field cricket Gryllus bimaculatus. Insect Mol Biol 2011, 20:619–635.
45. Yuan Q, Lin F, Zheng X, Sehgal A: Serotonin modulates circadian entrainment in Drosophila. Neuron 2005, 47:115–127.
46. Yuan Q, Joiner WJ, Sehgal A: A sleep-promoting role for the Drosophila serotonin receptor 1A. Can J Biochem 2006, 16:1051–1062.
47. Nichols CD: 5-HT2 receptors in Drosophila are expressed in the brain and modulate aspects of circadian behaviors. Dev Neurobiol 2007, 67:752–763.
48. Johnson O, Beitel C, Nichols CD: Serotonin 5-HT2 and 5-HT1A-like receptors differentially modulate aggressive behaviors in Drosophila melanogaster. Neuroscience 2009, 158:1292–1300.
49. Shimada M: Comparison of the Density-Dependent Population Process Between Laboratory and Wild Strains of Callosobruchus chinensis (L.) in Bruchids and Legumes: Economics, Ecology and Coevolution. Edited by Fujii K, Gatehouse AMR, Johnson CD, Mitchell R, Yoshida T. Dordrecht: Kluwer Academic; 1990:361–371.
50. Kondo N, Shimada M, Fukatsu T: High prevalence of Wolbachia in the azuki bean beetle Callosobruchus chinensis (Coleoptera, Bruchidae). Zool Sci 1999, 16:955–962.
51. Harano T, Miyatake T: Heritable variation in polyandry in Callosobruchus chinensis. Anim Behav 2005, 70:299–304.
52. Yasugi S, Miyatake T: Costs of mating and egg production in female Callosobruchus chinensis. J Insect Physiol 2003, 49:623–627.
53. Benjamin Y, Hochberg Y: Controlling the false discovery rate: a practical and powerful approach to multiple testing. J Roy Statist Soc B 1995, 57:289–300.
54. Quinn G, Keough MJ: Experimental design and data analysis for biologist. Cambridge: Cambridge University Press; 2002.
55. SAS Institute: JMP7. Cary, NC: SAS Institute Inc; 2007.