Physiological requirements for 20-hydroxyecdysone-induced rectal sac distention in the pupa of the silkworm, Bombyx mori

Suzuki Takumi, Sakurai Sho, Iwami Masafumi

Journal of Insect Physiology

Volume 56, Number 6, Page 673-677

doi: 10.1016/j.jinsphys.2010.02.006
Physiological requirements for 20-hydroxyecdysone-induced rectal sac distention in the pupa of the silkworm, Bombyx mori

Takumi Suzuki, Sho Sakurai, and Masafumi Iwami *

Division of Life Sciences, Graduate School of Natural Science and Technology, Kanazawa University, Kakumacho, Kanazawa 920-1192, Japan

* Corresponding author: Masafumi Iwami, Division of Life Sciences, Graduate School of Natural Science and Technology, Kanazawa University, Kakumacho, Kanazawa 920-1192, Japan

Tel: +81 76 264 6251
Fax: +81 76 264 6215
E-mail: masafumi@kenroku.kanazawa-u.ac.jp
Abstract

Successful insect development is achieved via appropriate fluctuation of ecdysteroid levels. When an insect’s ecdysteroid level is disrupted, physiological and developmental defects occur. In the pupa of the silkworm, Bombyx mori, the rectal sac is an essential organ that operates as a repository for degraded ecdysteroids, and it can be distended by administration of 20-hydroxyecdysone (20E). Our previous study showed that rectal sac distention appears 4 days after 20E administration. Hemolymph ecdysteroid levels, however, decrease to lower level during this period. Thus, the timing of the rectal sac distention does not match with that of ecdysteroid elevation. Here, we examine how 20E induces rectal sac distention. A ligature experiment and ecdysteroid quantification showed that continuous 20E stimulation induces rectal sac distention. Thorax tissue contributed to the continuous 20E stimulation needed to induce distention. Ecdysteroid released from the thorax tissue may be converted to 20E by ecdysone 20-hydroxylase to produce continuous 20E stimulation. Thus, the ecdysone metabolic pathway plays a critical role in rectal sac distention.

Keywords: excretory system, ecdysteroid, pupa, metamorphosis, rectal sac
1. Introduction

Ecdysteroids play critical roles in the regulation of insect development, and their titers in hemolymph are inseparably connected to developmental events. The hemolymph ecdysteroid titer is controlled by four steps: ecdysteroidogenesis in and release from the prothoracic glands, ecdysteroid uptake into target cells, ecdysteroid degradation at midgut, and excretion of the degraded ecdysteroid through the alimentary canal. Ecdysteroidogenesis in the prothoracic glands is activated by the prothoracicotropic hormone produced in the brain (Gilbert, 2004). When the brain is removed from the silkworm, *Bombyx mori*, the developmental sequence is arrested (Ichikawa and Ishizaki, 1961). In brain-removed pupae, the hemolymph ecdysteroid titer does not elevate and remains constant at a low level (Suzuki et al., 2009).

Insects show developmental defects when the ecdysteroid titer is disrupted. The larvae of *B. mori* fail to pupate when the ecdysteroid titer is decreased by ligature between the thorax and abdomen (Fukuda, 1944). By contrast, α-ecdysone, 20-hydroxyecdysone (20E), and inokosterone, a phytoecdysteroid, promote larval-pupal intermediates at high concentrations (Williams, 1968). A non-decreasing 20E titer inhibits both larval-pupal ecdysis and adult eclosion in the mealworm, *Tenebrio molitor* (Sláma, 1980) and the tobacco hornworm,
Manduca sexta (Truman et al., 1983). The inhibition may be due in part to the suppression of eclosion hormone release (Schwartz and Truman, 1983) and the consequent ecdysis triggering hormone release from Inka cells (Kigan and Adams, 2000). A decrease of the ecdysteroid titer is therefore necessary for successful ecdysis, adult eclosion, and further adult development (Schwartz and Truman, 1982); this is achieved by the degradation and excretion of ecdysteroid at each specific developmental stage.

Degraded ecdysteroid is transiently deposited in the rectal sac as meconium (Thompson et al., 1974). In B. mori, distention of the rectal sac occurs 4–5 days after pupation (Suzuki et al., 2009). Although distention can be induced by an injection of 20E, the ecdysteroid titer gradually decreases to a basal level in brain-removed pupae. Distention appears when the ecdysteroid titer decreases. Thus, the timing of rectal sac distention does not match that of ecdysteroid elevation. In the present study, we examined the physiological requirements for 20E-induced rectal sac distention. Our results indicate that tissue in the thorax is essential for 20E-induced rectal sac distention because it causes continuous 20E stimulation. In addition, they suggest that 20-hydroxylation of α-ecdysone also contributes to continuous 20E stimulation.
2. Materials and methods

2.1 Animals

*B. mori* (Kinshu × Showa) larvae were reared on an artificial diet (Silkmate 2M, Nihon Nosan Kougyo, Yokohama, Japan) at 25 ± 1°C under a 12 h light:12 h dark photoperiod. For our purposes, larvae a day before pupation are called prepupae. The day of pupation is designated day 0 (P0). One day after pupation and 2–8 days after pupation are designated stages P1 and P2–P8, respectively. Also, pupae just after larval-pupal ecdysis are called white pupae (WP).

2.2 Hormones and chemicals

α-Ecdysone and 20E were obtained from Sigma (St Louis, MO) and dissolved in ethanol and water, respectively. \[^{[3]}\text{H}\text{-Ecdysone (Perkin Elmer, Boston, MA)}\] was dissolved in borate buffer (100 mM boric acid, 50 mM borax, 60 mM NaCl). RH-5992, a non-steroidal ecdysone agonist (the kind gift of Y. Nakagawa of Kyoto University) was dissolved in dimethyl sulfoxide (DMSO). 20E, RH-5992, and DMSO were diluted with insect Ringer’s solution (128...
mM NaCl, 4.7 mM KCl, 1.9 mM CaCl₂) for injections. Each chemical, except for α-ecdysone and [³H]-ecdysone, was injected with various dosages per gram body weight.

2.3 Operation, hormonal treatment, and observations

Prepupae and WP pupae were ligated between head and thorax or thorax and abdomen by cotton thread followed by excision of the anterior remainder (referred to as isolated abdomens in the text). Two days after pupation, the isolated abdomens were injected with 3.0 µg/g 20E solution, 3.0 µg/g RH-5992 solution, or DMSO. Treated pupae were dissected at P8. Brain removal from WP pupa was performed as previously described (Suzuki et al., 2009). The pupae with brain-removed at WP were injected with 3.0 µg/g RH-5992 solution, DMSO, or the same volume of insect Ringer’s solution. The wound made by the operation or injection was sealed with melted paraffin wax. Rectal sac distention was observed as previously described (Suzuki et al., 2009).

2.4. Quantification of ecdysteroid titer

Hemolymph was collected from the pupae through an incision on the dorsal side.
Ecdysteroids were extracted from the hemolymph and quantified by radioimmunoassay as previously described (Sakurai et al., 1998). Anti-ecdysone antiserum H-22 was obtained from L. I. Gilbert and D. H. S. Horn and used as a capture antibody in the radioimmunoassay (Warren and Gilbert, 1986). We examined the cross-reactivity of anti-ecdysone antiserum H-22 against RH-5992; no cross-reactivity was found (data not shown).

2.5. Reverse transcription (RT)-PCR

Total RNA was extracted from pupal midgut by Trizol reagent (Invitrogen) and then treated with DNase I (Promega, Madison, WI). Complementary DNA was prepared from 0.6 µg of total RNA with reverse transcriptase ReverTra Ace (Toyobo, Osaka, Japan) by an anchored oligo-dT according to the manufacture’s protocol. The PCR primer were designed according to the nucleotide sequences: ribosomal protein L3 (RpL3), 5’-AGCACCCCCGTGCTGCTGCTGCTA-3’ and 5’-TGGGTGTTGTGGGAGGTCTG-3'; ecdysone receptor isoform A (EcR-A), 5’-TGCGTCCAAGCTCATCCTGC-3’; ecdysone receptor isoform B1 (EcR-B1), 5’-AGCAGGGTTGCGGTTTCCCGCTGCG-3’ and 5’-CGGTGTGTTGAGGACATTGGA-3’; ultraspiracle l (Usp-l),
5’-GTCGAGCGTGCGAAGAAA-3’ and 5’-CAGCCATTGTATATCGAGTTCA-3’;

Ultraspiracle 2 (Usp-2), 5’-GATATCGTGATAATAACCTAAGTA-3’ and 5’-GCAACAAGGTCGTGGAACTAA-3’; shade, 5’-TTGTAGCTGTAACCGCGTTG-3’ and 5’-GTACGGCCTCAAGAGCAGTC-3’. These primers were designed according to previous report (Kaneko et al., 2006) except primers for shade (Maeda et al., 2008). PCR was carried out with Go Taq (Promega) under following conditions: denature at 94 °C for 30 seconds, annealing at different temperatures for each gene (RpL3, EcR-A, and EcR-B1: 60°C; Usp-1: 62°C; Usp-2: 56°C; shade: 57°C) for 30 seconds, and extension at 72 °C for 30 seconds, followed by 1 minute as a final extension. RpL3 was used as an internal standard and amplified for 30 cycles. EcR-A, EcR-B1, Usp-1, Usp-2, and shade were amplified for 35, 35, 37, 37, and 33 cycles, respectively. The products were cloned into pGEM-T vector (Promega) and subjected to sequence analysis. The sequences of these plasmids were expected ones, indicating that each sequence was correctly amplified under the present experimental condition.

3. Results

3.1 The thorax was required for rectal sac distention induced by 20E
Distention appears 4 days after 20E injection in brain-removed pupae (Suzuki et al., 2009). When we monitored the ecdysteroid titer, by the fourth day it had fallen below 3.0 µM (Fig. 1). Although rectal sac distention is induced by 20E (Suzuki et al., 2009), the timing of successful distention does not coincide with that of ecdysteroid elevation. There are two possible explanations for this time lag: either 20E elevation triggers the initiation of distention, but distention requires 4 days to take place, or the ecdysteroid decrease that follows the elevation induces distention. To distinguish between these hypotheses, we prepared two different pupae: brain-removed and ligated pupae. WP pupae were either brain-removed or ligated between thorax and abdomen (isolated abdomen), injected with 3.0 µg/g 20E or insect Ringer’s solution, and dissected at P6. An injection of 20E successfully induced distention in 87% of the brain-removed pupae and in 20% of the isolated abdomens (Table 1). In the control experiment, insect Ringer’s solution did not induce distention.

We then used pupae ligated a day before pupation. Prepupae were ligated between head and thorax (decapitated) or between thorax and abdomen, then injected with 3.0 µg/g 20E or insect Ringer’s solution and dissected at P6 and P8. At P6 and P8, 24% and 53%, respectively, of the decapitated pupae injected with 20E exhibited distention (Table 2). Distention did not
appear in the decapitated pupae injected with insect Ringer’s solution. The isolated abdomens injected with 20E did not show distention at either P6 or P8. Thus, the decapitated pupae exhibited distention as the result of an injection of 20E, but the isolated abdomens did not exhibit any, indicating that the thorax was necessary to successfully achieve distention.

3.2 Continuous stimulation of 20E was required for successful distention

Injection of 20E restored rectal sac distention in the decapitated pupae but not in the isolated abdomens (Table 2). Decapitated pupae contain a larger amount of ecdysteroid than isolated abdomens because the prothoracic glands produce ecdysteroid in the pupae, indicating that successful distention requires continuous exposure to ecdysteroid. Isolated abdomens do not contain sufficient endogenous ecdysteroid probably because they lack prothoracic glands and cannot achieve successful development without exogenous ecdysteroid. 20E may expire easily, since injected 20E has a half-life as short as 4 hours (Nijhout, 1976). We therefore attempted to give isolated abdomens multiple injections of 20E. Abdomens isolated at the prepupal stage were injected with 0.25–0.5 µg/g 20E during P2–P4 and dissected at P8. Double injections of 0.5 µg/g 20E at P2 and P3 and triple injections of 0.25 µg/g 20E every P2–P4 did
not restore distention (Table 3). By contrast, triple injections of 0.5 µg/g 20E significantly
restored distention (97%), indicating that induction of distention in isolated abdomens requires
continuous stimulation by 20E.

3.3 The thorax was necessary to keep the ecdysteroid level high enough to induce distention

Table 3 indicates that decapitated pupae required a 20E injection, but isolated abdomens required triple injections for successful rectal sac distention. These results imply that injected 20E was degraded in the isolated abdomens. We quantified the hemolymph ecdysteroid titer in both decapitated pupae and isolated abdomens injected with 20E. The decapitated pupae and isolated abdomens were injected with 3.0 µg/g of 20E or insect Ringer’s solution at P2, and hemolymph was collected during P3–P8. The ecdysteroid titers during P1–P2 were 1.11 ± 0.26 – 1.26 ± 0.20 µM and 0.45 ± 0.08 – 0.92 ± 0.20 µM in the decapitated pupae and isolated abdomens, respectively (Fig. 2). After an injection with 20E, ecdysteroid titers were significantly elevated in both the decapitated pupae and isolated abdomens, and they remained at 4.4–5.2 µM and 3.2–3.7 µM, respectively. Insect Ringer’s solution did not cause any change in the ecdysteroid titer. Thus, the hemolymph ecdysteroid titer in the decapitated pupae
remained at a higher level than that in the isolated abdomens. This higher ecdysteroid level may
be sufficient for achievement of continuous stimulation of 20E in the decapitated pupae.

3.4 Rectal sac distention was induced by non-degradative ecdysone

We confirmed that continuous stimulation by 20E induces rectal sac distention in the
decapitated pupae by using a non-degradative ecdysone agonist, RH-5992. RH-5992 has a
longer half-life than 20E (Retnakaran et al., 1995) and did not affect de novo ecdysteroid
synthesis (Fig. 3). The effects of RH-5992 on rectal sac distention can be examined without
interference from endogenous ecdysteroid. Hemolymph was collected from pupae injected with
RH-5992 or DMSO at P4 and P5. There was no statistically significant difference between the
ecdysteroid titer of pupae injected with RH-5992 and that of the pupae injected with DMSO at
either stage. Pupae were brain-removed at the WP stage, injected with 3.0 µg/g RH-5992 at P2,
and dissected at P8. As a control, insect Ringer’s solution and DMSO were also injected. After
an injection of RH-5992, 75% of the pupae showed successful distention (Table 4). No pupae
exhibited distention as a result of injection with insect Ringer’s solution or DMSO.

In isolated abdomens, successful distention was only achieved by triple injections of
20E (Table 3). A single injection of RH-5992 induced distention in isolated abdomens. RH-5992
injected at P2 induced rectal sac distention in 58% of the isolated abdomens by P8 (Table 5).
DMSO did not induce distention in the isolated abdomens. Thus, a single injection of
non-degradative ecdysone agonist induced rectal sac distention in isolated abdomens, indicating
that successful distention requires continuous ecdysteroid stimulation and does not require a
decrease in ecdysteroid titer.

3.5 Ecdysone receptor genes and ecdysone 20-hydroxylase gene were expressed in the midgut

We considered the reason why single injection of 20E was able to induce rectal sac
distention in the decapitated pupae. An injection of 20E stimulates ecdysone 20-hydroxylase
activity in the midgut of M. sexta (Keogh et al., 1989) and Spodoptera littoralis (Williams et al.,
1997). We therefore examined 20E effects on expression of ecdysone 20-hydroxylase gene in
the pupal midgut. First, expression levels of ecdysone receptors were examined by
semi-quantitative RT-PCR to confirm that 20E acted in the midgut. The expressions of EcR-A
and EcR-B1 were kept at constant levels throughout P0 – P6 (Fig. 4). Similarly, usp-1 and usp-2
were also expressed at constant levels during P0 – P6. Throughout the periods, 20E was able to
act in the midgut since ecdysone receptor genes were expressed in the tissue constantly. Second, we examined the expression profile of ecdysone 20-hydroxylase gene, *shade* (Petryk *et al.*, 2003; Maeda *et al.*, 2008) and found that the level of *shade* expression in the midgut during P0 – P6. The gene *shade* was thus expressed at constant level during examined periods (Fig.4).

4. Discussion

In this study, we explored the physiological requirements for 20E-induced rectal sac distention, and we found that thorax tissue is required. An injection of 3.0 µg/g 20E induced rectal sac distention in over 50% of decapitated pupae, but it did not induce distention in isolated abdomens (Table 3), indicating that the thorax is necessary for rectal sac distention. Triple injections of over 0.5 µg/g 20E or a single injection of RH-5992 successfully induced distention in isolated abdomens (Tables 3, 5). This result indicates that continuous ecdysteroid stimulation is necessary for distention to occur. In decapitated pupae, continuous 20E stimulation can be achieved by a single injection, while a triple injection of 20E or an injection of RH-5992 can induce distention in isolated abdomens. For continuous ecdysteroid stimulation, injected 20E could stimulate ecdysteroid synthesis in thorax tissue. Exogenous 20E stimulates
ecdysteroid synthesis in the prothoracic glands in the pupae of *M. sexta* (Sakurai and Williams, 1989). Regardless of the presence of prothoracic glands, an injection of 20E induced rectal sac distention in 87% of brain-removed pupae (Table 1) and in only 24% of decapitated pupae at P6 (Table 3). The different responsiveness may be due to a change of 20E action to prothoracic glands during metamorphosis (Sakurai and Williams, 1989).

The gene encoding ecdysone 20-hydroxylase, enzyme that promotes conversion of \( \alpha \)-ecdysone to 20E (Gilbert, 2004), was expressed constantly in the midgut during P0 – P6 (Fig. 4). Ecdysteroid from the prothoracic glands may contribute to inducing rectal sac distention by acting as a source of 20E, since an injection of 20E induced rectal sac distention in the decapitated pupae but did not in the isolated abdomens. In fact, the ecdysteroid titer in decapitated pupae was almost 2-fold higher than that in isolated abdomens (Fig. 2). Thus, endogenous \( \alpha \)-ecdysone may contribute to continuous 20E stimulation to successfully induce distention.

Injected 20E degenerates rapidly in the larvae of *M. sexta*, and the half-life of 20E is approximately 4 hours in isolated abdomens (Nijhout, 1976). We previously reported that the ecdysteroid titer decreased to a basal level 4 days after injection of brain-removed pupae with
20E (Suzuki et al., 2009). However, in decapitated pupae and in isolated abdomens, hemolymph ecdysteroid titers remained at constant levels after 20E injection (Fig. 2). The ecdysteroid titers were quantified by radioimmunoassay using anti-ecdysone antiserum as a capture antibody. The antiserum recognizes both active and inactive ecdysteroid (Warren and Gilbert, 1986). The detected ecdysteroids could be a mixture of active and inactive ecdysteroids because rectal sac distention required continuous 20E stimulation and distention was not induced in isolated abdomens (Table 2, 3). Thus, 20-hydroxylation may be required for continuous stimulation to successfully induce distention. Although it is not clear what causes the different ecdysteroid titers between brain-removed pupae and ligated pupae after 20E injection, our results suggest that the ecdysteroid metabolic pathway is involved in the sequence of 20E-induced rectal sac distention.

Acknowledgements

We are grateful to Drs. L. I. Gilbert and D. H. S. Horn for the anti-ecdysone antiserum H-22, Dr. Y. Nakagawa for the RH-5992, Dr. Manaporn Manaboon and Masami Miyakoshi of our laboratory for collecting the white pupae, and Yuichiro Nagamura of
Radioisotope Laboratory for Natural Science and Technology, Kanazawa University, for technical advice. This work was supported in part by Grants-in-Aid for Scientific Research (18380040) and Exploratory Research (19658019) from the Japan Society for the Promotion of Science.

References

Fukuda, S., 1944. The hormonal mechanism of larval molting and metamorphosis in the silkworm. Journal of Faculty of Science Tokyo Imperial University 4, 477–532.


Kaneko, Y., Takaki, T., Iwami, M., Sakurai, S., 2006. Developmental profile of Annexin IX and


1 1,2-dibenzoyl-1-tert-butylhydrazine (RH-5849). Journal of Biological Chemistry 272, 8427-8432.

3
Figure legends

Fig. 1. Rectal sac distention and hemolymph ecdysteroid titer. (A) Pupae were dissected between P3 and P6. Rectal sacs distended between P5 and P6, and the ecdysteroid titer (open circle) started to decrease at P5. (B) Pupae were brain-removed at WP stage and injected with 3.0 µg/g 20E solution 2 days after brain removal. Sacs distended 3–4 days after injection, and the titer (open circle) started to decrease 3 days after injection. Successful distention is expressed as the ratio of the number of pupae that show distended sacs to that of total pupae. The arrow indicates the day of 20E injection. Each datum indicates the mean ± standard deviation. Data are taken and modified from Suzuki et al., 2009.

Fig. 2. Ecdysteroid titers in normal pupae, decapitated pupae, and isolated abdomens. Prepupae were ligated between head and thorax (decapitated pupae, closed circles) or thorax and abdomen (isolated abdomens, triangles) and injected with 3.0 µg/g of 20E solution 2 days after pupation. As a control (open circles), decapitated pupae were injected with insect Ringer’s solution at P2. The titers in normal pupae are indicated as open squares. Each datum indicates the mean ± standard deviation ($n = 6 – 10$).
Fig. 3. Ecdysteroid titers in brain-removed pupae injected with RH-5992. Pupae were brain-removed at WP stage and injected with 3.0 µg of RH-5992 (closed bars) or DMSO (open bars) 2 days after brain removal. Hemolymph was collected from the pupae 4 and 5 days after pupation.

The ecdysteroid titer was quantified by radioimmunoassay. The concentration of ecdysteroid is presented as α-ecdysone equivalent. Each datum indicates the mean ± standard deviation (n = 8).

Fig. 4. Temporal expression profiles of EcR-A, EcR-B1, Usp-1, Usp-2, shade, and RpL3 in the midgut during P0 – P6. The gene RpL3 was used as internal standard. The term –RT indicates that the RNA samples were subjected to PCR without reverse transcriptional reaction. The expression profiles were analyzed 3 times by independent experiments and show no difference in the experiments.
Table 1 20E did not induce rectal sac distention in isolated abdomens of white pupae

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Chemical</th>
<th>Total</th>
<th>Distention</th>
<th>No distention</th>
<th>Dead</th>
<th>Ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain removal</td>
<td>Ringer’s</td>
<td>15</td>
<td>0</td>
<td>15</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>20E</td>
<td>15</td>
<td>13</td>
<td>2</td>
<td>0</td>
<td>87</td>
</tr>
<tr>
<td>Abdominal isolation</td>
<td>Ringer’s</td>
<td>13</td>
<td>0</td>
<td>13</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>20E</td>
<td>15</td>
<td>3</td>
<td>12</td>
<td>0</td>
<td>20</td>
</tr>
</tbody>
</table>

All numerals indicate the number of pupae except ratios, which are expressed as percentages.
Table 2  20E induced rectal sac distention in decapitated pupae but not in isolated abdomens treated 1 day before pupation

<table>
<thead>
<tr>
<th>Stage</th>
<th>Treatment</th>
<th>Total</th>
<th>Distention</th>
<th>No distention</th>
<th>Dead</th>
<th>Ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P6</td>
<td>Intact</td>
<td>21</td>
<td>21</td>
<td>0</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Decapitated + Ringer’s</td>
<td>16</td>
<td>0</td>
<td>16</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Decapitated + 20E</td>
<td>29</td>
<td>7</td>
<td>20</td>
<td>2</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>Isolated abdomen + 20E</td>
<td>32</td>
<td>0</td>
<td>29</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>P8</td>
<td>Decapitated + Ringer’s</td>
<td>16</td>
<td>1</td>
<td>13</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Decapitated + 20E</td>
<td>30</td>
<td>16</td>
<td>14</td>
<td>0</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>Isolated abdomen + 20E</td>
<td>31</td>
<td>3</td>
<td>25</td>
<td>3</td>
<td>10</td>
</tr>
</tbody>
</table>

All numerals indicate the number of pupae except ratios, which are expressed as percentages.
Table 3 Multiple injections of 20E restored rectal sac distention in isolated abdomens

<table>
<thead>
<tr>
<th>Dose</th>
<th>Total</th>
<th>Injections</th>
<th>Distention</th>
<th>No distention</th>
<th>Dead</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25 µg</td>
<td>30</td>
<td>2</td>
<td>4</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>0.5 µg</td>
<td>38</td>
<td>2</td>
<td>4</td>
<td>33</td>
<td>1</td>
</tr>
<tr>
<td>0.5 µg</td>
<td>35</td>
<td>3</td>
<td>34</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

All numerals indicate the number of pupae except the 20E dosage and the number of injections.
Table 4 RH-5992 induced rectal sac distention in brain-removed pupae

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Total</th>
<th>Distention</th>
<th>No distention</th>
<th>Ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ringer’s</td>
<td>10</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>DMSO</td>
<td>12</td>
<td>0</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>RH-5992</td>
<td>12</td>
<td>9</td>
<td>3</td>
<td>75</td>
</tr>
</tbody>
</table>

All numerals indicate the number of pupae except ratios, which are expressed as percentages.
Table 5 RH-5992 induced rectal sac distention in isolated abdomens

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Total</th>
<th>Distention</th>
<th>No distention</th>
<th>Dead</th>
<th>Ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20E</td>
<td>15</td>
<td>0</td>
<td>15</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>DMSO</td>
<td>26</td>
<td>0</td>
<td>26</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>RH-5992</td>
<td>31</td>
<td>18</td>
<td>12</td>
<td>1</td>
<td>58</td>
</tr>
</tbody>
</table>

All numerals indicate the number of pupae except ratios, which are expressed as percentages.
The occurrence of distention (%)

Ecdysteroid (µM)

Days after pupation

(A)

Days after injection

(B)