

# Comparisons in Temperature and Photoperiodic-dependent Diapause Induction between Domestic and Wild Mulberry Silkworms

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## Research Article

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**Comparisons in temperature and photoperiodic-dependent diapause induction  
between domestic and wild mulberry silkworms**

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## Abstract

The bivoltine strain of the domestic silkworm, *Bombyx mori*, has two generations per year. It shows a facultative diapause phenotype determined by environmental conditions, including photoperiod and temperature, and nutrient conditions during embryonic and larval development of the mother. However, it remains unclear how the environmental signals received during development are selectively utilized as cues to determine alternative diapause phenotypes. We performed a comparative analysis between the Kosetsu strain of *B. mori* and a Japanese population of the wild mulberry silkworm *B. mandarina* concerning the hierarchical molecular mechanisms in diapause induction. Our results showed that for the Kosetsu, temperature signals during the mother's embryonic development predominantly affected diapause determination through the thermosensitive transient receptor potential ankyrin 1 (TRPA1) and diapause hormone (DH) signaling pathways. However, embryonic diapause in *B. mandarina* was photoperiod-dependent, although the DH signaling pathway and thermal sensitivity of TRPA1 were conserved within both species. Based on these findings, we hypothesize that TRPA1-activated signals are strongly linked to the signaling pathway participating in diapause induction in Kosetsu to selectively utilize the temperature information as the cue because temperature-dependent induction was replaced by photoperiodic induction in the TRPA1 knockout mutant.

## Introduction

In the domestic silkworm *Bombyx mori*, progeny diapause is induced during the early embryonic stage. The development of diapause-destined embryos is arrested during the G2 cell cycle stage immediately after forming the cephalic lobe and telson and sequential segmentation of the mesoderm and ectoderm<sup>1</sup>. In bivoltine strains, populations of *B. mori* that have two generations per year show a facultative diapause phenotype, which is determined by environmental conditions, such as photoperiod, temperature, and nutrient conditions during embryonic and larval development in the mother's generation<sup>2</sup>. In general, progeny diapause of bivoltine strains is predominantly determined by environmental conditions during maternal embryonic development in conjunction with long days and high temperatures that induce diapause eggs<sup>2-4</sup>. However, the environmental cues that induce diapause are different in each bivoltine strain and are also updated by integrating cumulative signals received through development<sup>5,6</sup>. For instance, when larvae were reared on an artificial diet under long-day conditions, some silkworm strains produced non-diapause eggs even if the eggs were incubated in diapause egg-producing conditions<sup>2,7</sup>. However, it remains unclear how environmental signals received through development are selectively utilized as cues to determine alternative diapause phenotypes.

For the bivoltine strain Kosetsu, the temperature signal during the mother's embryonic development is the predominant factor affecting diapause determination, regardless of the photoperiods during embryonic and larval development under our rearing condition using an artificial diet<sup>8,9</sup>. When incubating eggs at 25°C under continuous darkness (25DD), the resultant female moths lay nearly 100% diapause eggs. In contrast, incubation of eggs at 15°C in dark conditions (15DD) resulted in moths that

lay nearly 100% non-diapause eggs. Thus, Kosetsu is considered a typical strain that is susceptible to temperature-dependent diapause induction.

The molecular mechanisms underlying the induction of *B. mori* embryonic diapause have been investigated extensively<sup>5,9</sup>. Our previous study revealed that embryonic diapause is induced by the diapause hormone (DH) signaling pathway, consisting of a highly sensitive and specific interaction between DH and DH receptors (DHR) pupal-adult development<sup>8</sup>. *DH-PBAN* encodes a polyprotein precursor containing DH, pheromone biosynthesis activating neuropeptide (PBAN), and  $\alpha$ -,  $\beta$ -, and  $\gamma$ -subesophageal ganglion (SG) neuropeptides (SGNPs). DH is produced from the DH-PBAN precursor through posttranslational processing<sup>10</sup>. It is produced exclusively in seven pairs of neurosecretory cells, known as DH-PBAN-producing neurosecretory cells (DHPCs), located within the SG in the mother's generation<sup>11</sup>. DH is released from the corpus cardiacum (CC)<sup>12</sup>, a major release site for neuropeptide hormones, into the hemolymph, and acts on DHR in ovary<sup>13</sup>. Furthermore, the cerebral  $\gamma$ -aminobutyric acid (GABA)ergic and corazonin (Crz) pathway modulates DH release throughout the temperature-dependent expression of the plasma membrane GABA transporter<sup>2</sup>. Recent studies revealed that the transient receptor potential ankyrin 1 (TRPA1) (BmoTRPA1) acts as a thermosensitive channel activated at temperatures above  $\sim 21^{\circ}\text{C}$  and affects diapause induction through DH release in Kosetsu. Thus, BmoTRPA1 acts as a molecular switch for the temperature-dependent diapause induction in Kosetsu<sup>14</sup>.

*B. mori* was domesticated from the wild mulberry silkworm, *B. mandarina*, approximately 5,000–10,000 years ago in China<sup>15–18</sup>. *B. mori* and *B. mandarina* are similar in morphological, cytological, and physiological aspects; copulation between them is possible, and the resultant progeny is never sterile<sup>19–21</sup>, although both species are

clearly genetically differentiated<sup>22</sup>. In addition, *B. mandarina* enters embryonic diapause at a similar developmental stage as in *B. mori*<sup>23,24</sup>. The bi-, tri-, and univoltine *B. mandarina* inhabit Japan based on the moth's seasonal fluctuation<sup>25,26</sup>. Furthermore, photoperiodic-dependent diapause induction of *B. mandarina* was shown in a previous report, in which more female moths produced diapause eggs under short-day conditions, such as a 12-h light/12-h dark cycle (12L12D) than under long-day conditions, such as 15L9D during larval stages, even if the temperature was kept constant at 25°C during embryonic development. Diapause induction is most strongly affected by photoperiod in a certain developmental stage following the second larval ecdysis<sup>27</sup>. Thus, to our knowledge, *B. mandarina* enters diapause by photoperiodic induction, which contrasts with the temperature-dependent diapause induction in Kosetsu of *B. mori*. Therefore, we performed a comparative analysis of diapause induction using Kosetsu and a Japanese population of *B. mandarina* with respect to the hierarchical molecular mechanisms to elucidate the selectivity and integration of environmental cues. In the present study, we hypothesize that TRPA1-mediated signals are strongly linked to the signaling pathway participating in diapause induction in Kosetsu to selectively utilize the temperature information as the cue.

## Results and discussion

### Temperature and photoperiodic-dependent diapause induction are different between the Kosetsu strain of *B. mori* and the Fuchu population of *B. mandarina*.

We first confirmed the photoperiodic response in diapause induction of *B. mandarina* as described previously<sup>27</sup>. When eggs were incubated under 25DD during embryonic development, and thereafter, the larvae reared under long-day conditions,

16L8D and 24L0D, eclosed moth laid non-diapause eggs that hatched 10 d after oviposition (Figs. 1a, b). In contrast, most eggs were pigmented and never hatched 30 d after oviposition under short-day condition, 8L16D, for 25DD embryos. These pigmented eggs arrested normal embryogenesis, and the arrested stage was similar to that in diapause eggs in *B. mori* (Figs. 1a, c). The progeny eggs entered diapause under all photoperiodic conditions when embryos were reared under 25DD in *B. mori* (Fig. 1b), indicating that there were differences in the environmental cues also temperature or photoperiodic-dependent diapause induction in both strains. However, both strains never entered diapause for progeny from the 15DD condition (Fig. 1b).

#### **DH induces progeny diapause in *B. mandarina*.**

To clarify whether the DH signaling pathway is involved in diapause induction in *B. mandarina*, we determined the complementary DNA (cDNA) sequences of *DH-PBAN* and *DHR*. The *B. mandarina DH-PBAN* cDNA was cloned, and the deduced amino acid sequence of DH (BmaDH) was identical to that of *B. mori* DH (BmoDH)<sup>28</sup> (Supplementary Fig. S1). We also found the homologous sequence to *B. mori DHR* (*BmoDHR*) cDNA using SilkBase (<http://silkbases.ab.a.u-tokyo.ac.jp/cgi-bin/index.cgi>). Using reverse transcription PCR (RT-PCR), we cloned *B. mandarina DHR* cDNA (*BmaDHR*; Acc. No. LC594680). The sequence of the deduced amino acids in BmaDHR represented a highly similar sequence in BmoDHR at 97%, including the seven transmembrane regions (TM1–TM7) (Supplementary Fig. S1). Next, immunostaining was performed using an anti-DH antibody. High-immunofluorescent signals were detected as a cluster of somata in SG (Figs. 2a,c), considering that these immunoreactive clusters of somata were identical to SMd, SMx, SLb, and SL neurons of DHPCs in *B.*



*mori*<sup>12</sup> (Figs. 2b,c). Furthermore, immunoreactive neurites with bead-like varicosities were detected in the CC and projected onto the corpus allatum (CA) surface to form blind ends like that of *B. mori* (Fig. 2d), suggesting that BmaDH was released to the hemolymph through these neurohemal organs.

Next, we injected synthetic DH into *B. mandarina* pupae developed from 25DD and 15DD embryos and reared under the 16L8D condition (Figs. 2e). Although peanut oil-injected silkworms never oviposited diapause eggs, DH-injected silkworms oviposited diapause eggs in a dose-dependent manner. The diapause eggs were staged in the early embryonic stages, which was similar to the cases of diapause eggs of *B. mandarina* (8L16D) and *B. mori* (Fig. 1c, 16L8D\_+DH). Therefore, we concluded that the DH signaling pathway is conserved between *B. mori* and *B. mandarina* to induce progeny diapause.

### **Cloning and functional analysis of TRPA1 orthologs in *B. mandarina*.**

We cloned two TRPA1 cDNAs, namely *BmaTRPA1a* (Acc. No. LC597009.1) and *BmaTRPA1b* (Acc. No. LC597010.1), from *B. mandarina* eggs just before hatching. The amino acid sequences deduced from the two cDNAs (*BmaTRPA1a* and *BmaTRPA1b*) were highly similar to that of BmoTRPA1 at 99% and 97%, respectively (Fig. 3a and Supplementary Fig. S2). The *BmaTRPA1* genes are conserved in the functional domain consisting of ankyrin repeat domains (ANK1~17), transmembrane domains (TM1~6), and a pore-loop domain among the TRPA1 subfamily<sup>29,30</sup>. *BmaTRPA1a* and *BmaTRPA1b* transcripts were caused by alternative splicing of distinct exons encoding the linker region located between the last ankyrin repeat and the first transmembrane segment (Figs. 3a, 3b, and Supplementary Fig. S2). This structure

resembles the alternative splicing variants of TRPA1 found in *Drosophila melanogaster* and *Aedes aegypti*<sup>29,30</sup>.

Next, channel properties of TRPA1 between the two species were compared by an electrophysiological approach. BmaTRPA1 or BmoTRPA1 were heterologously expressed in *Xenopus* oocytes, and ionic currents were observed. TRPA1 from both species was activated by heat. We investigated temperature sensitivity and determined the threshold temperature for activation for BmaTRPA1a and BmaTRPA1b as well as BmoTRPA1 with an Arrhenius plot (Fig. 3c–f). The average thermal activation threshold of BmoTRPA1 was  $24.0^{\circ}\text{C} \pm 0.8^{\circ}\text{C}$  (Fig. 3c). This value is slightly higher than that of BmoTRPA1 previously reported using mammalian cultured cells (approximately  $21^{\circ}\text{C}$ )<sup>14</sup>, which might be caused by different cell types used for heterologous expression of TRPA1. The average thermal activation thresholds of BmaTRPA1a and BmaTRPA1b were  $24.4^{\circ}\text{C} \pm 0.6^{\circ}\text{C}$  and  $24.0^{\circ}\text{C} \pm 0.4^{\circ}\text{C}$ , respectively. No statistical difference was observed among thermal activation thresholds for BmoTRPA1, BmaTRPA1a, and BmaTRPA1b (Fig. 3f). Therefore, we concluded that the embryonic expression and temperature sensitivity of TRPA1 are conserved in both species.

#### **Knockout (KO) analysis of BmoTRPA1 in diapause induction.**

Since both temperature-sensitive TRPA1 and DH signaling pathways are conserved in *B. mori* and *B. mandarina*, we hypothesized that there were differences in the downstream signal of TRPA1 activation to induce diapause between *B. mori* and *B. mandarina*. Hence, we constructed a KO mutant of *BmoTRPA1* to investigate the TRPA1-activated downstream pathway. We designed a transcription activator-like effector nuclease (TALEN) target in the second exon encoding the first ankyrin repeat

(ANK1) (Figs. 3a, 4a, and Supplementary Fig. S2). We isolated a homozygous mutant in which both a 7-base sequence was deleted, and a 9-base sequence was inserted into the spacer region between two TALEN binding sites, and a single base pair was replaced in the right TALEN binding site (TAL2) and designated the mutant as  $\Delta TRPA1\_1429$  (Figs. 4a, b).  $\Delta TRPA1\_1429$  was considered a null mutant, which could not translate full-length *BmoTRPA1* due to a frameshift of *BmoTRPA1* cDNA. We then investigated whether the  $\Delta TRPA1\_1429$  silkworm oviposited non-diapause eggs from 25DD and 15DD (Figs. 4c, d). Although the wild type (*wt*) moths laid 100% of diapause eggs, all  $\Delta TRPA1\_1429$  moths laid only non-diapause eggs when the larvae were reared under the long-day condition (16L8D). In this condition, the hemolymph DH levels of  $\Delta TRPA1\_1429$  from 25DD embryos were extremely low compared with those of *wt* for both samples collected 2 and 4 days after pupation, resulting in decreased levels similar to the 15DD of *wt* (Fig. 4e). Furthermore, when picrotoxin (PTX), Crz, or DH were injected into  $\Delta TRPA1\_1429$  pupa under the 16L8D condition, higher diapause egg-inducing activity was observed as that in 15DD silkworms (Fig. 4d), suggesting that the GABAergic, Crz, and DH signaling pathways were retained in  $\Delta TRPA1\_1429$ .

However, the ratios of non-diapause eggs gradually decreased depending on the length of scotophase in  $\Delta TRPA1\_1429$ . The percentages of diapause eggs were 14.7% and 84.6% in 13L11D and 8L16D, respectively (Fig. 4c). Taken together, we speculated that although the *BmoTRPA1*-activated pathway is involved in diapause induction in the *wt* of *Kosetsu* through the GABAergic, Crz, and DH signaling pathways, the disruption of the *TRPA1*-activated pathway in the *BmoTRPA1* KO mutant could not operate the temperature-dependent diapause induction. Interestingly, *BmoTRPA1* KO mutants' progeny eggs exhibited photoperiodic diapause induction, as is the case with *B.*

*mandarina*. This suggests that the *wt* of Kosetsu potentially possesses a signaling pathway involved in photoperiodic diapause induction that might be canceled by the strong linkage of the BmTRPA1-activated signal and DH signaling pathways. Consequently, the progeny diapause may be dependent on photo- or scotophase-length in *BmoTRPA1* KO mutants such as *B. mandarina*. To our knowledge, the temperature-dependent diapause induction did not occur in *B. mandarina*. Therefore, we speculated that temperature-dependent diapause induction spreads through the artificial selection process during domestication because temperature control by humans is easier than photoperiodic control by human handling. Furthermore, as it is suggested that insect clocks are engaged in photoperiodic response, including diapause induction<sup>31</sup>, both the temperature and photoperiodic diapause induction may be mediated by common unidentified machinery such as circadian clocks. In this study, we investigated to elucidate the molecular mechanism of selectivity and integration of environmental cues concerning diapause induction. Our results demonstrated the usefulness of a comparative study using the Kosetsu strain of *B. mori* and a Japanese population of *B. mandarina* for further research on diapause induction.

## **Methods**

### **Silkworms**

For *B. mori*, the bivoltine Kosetsu strain was used in these experiments. Eggs were incubated under two different conditions: 1) at 25°C under continuous darkness (25DD) to obtain diapause eggs in the *wt*, and 2) at 15°C under continuous darkness (15DD) to obtain non-diapause eggs. Larvae were then reared on an artificial diet (Kuwano-hana for young silkworm, Gunma Artificial Diet Production Center, Takasaki,

Gunma) at 25°C–27°C under 16-h light/8-h dark (16L8D), 13L11D, 8L16D, and 24L0D cycles at a relative humidity of 30%–50%. Pupae used in the experiments were collected within 1 h after each ecdysis (referred to as day 0) to synchronize their subsequent development. Pupae were incubated at 25°C to allow for adult development. The larvae of wild mulberry silkworm, *B. mandarina*, were collected from Tokyo University of Agriculture and Technology in Fuchu (35°41'N) and the Gunma Sericultural Technology Center at Maebashi (36°24'N). The oviposited non-diapause eggs were incubated at 25DD and 15DD. Larvae were reared on mulberry leaves at 25°C–27°C under 16L8D, 8L16D, and 24L0D. Pupae used in the experiments were collected within 12 h after each ecdysis. Pupae were incubated at 25°C to allow for adult development.

The percentage of diapause eggs was estimated by counting the number of eggs in diapause and those not in diapause in each egg batch after the non-diapause eggs hatched. The results are expressed as the average percentage of diapause in each egg batch<sup>8,9</sup>. For *B. mandarina*, the eggs were counted using a SZ61 stereo microscope (Olympus, Tokyo, Japan). Pupal-adult development lasted 12–26 d in these experiments, and a few pupae did not eclose within one month, as described in a previous report<sup>32</sup>. The unclosed pupae were not used for calculation of the diapause eggs inducing activity.

## **Thionin staining and immunostaining**

Thionin staining was performed as previously described<sup>8</sup>. The eggs were fixed in Carnoy's fixative for 1 d at 4°C. The fixed eggs were treated with a gradient concentration of ethanol for 10 min. Following 5 min of boiling, the chorion was dissected, and the eggs were transferred into a thionine solution (0.07% thionine and 0.3% phenol dissolved in 80% ethanol) for 2 h. The stained eggs were rinsed with 80%

ethanol four times and dehydrated with a gradient concentration of ethanol for 10 min after each rinse. The eggs were then soaked in benzene to make the yolk transparent. The embryos were observed using an M165 FC stereomicroscope (Leica, Wetzlar, Germany).

The immunoreaction procedures were adapted as previously reported<sup>11</sup>. The brain-SG complex or the CC and CA complex were dissected in fixative containing 4% paraformaldehyde, 7% picric acid, 10 mM MgCl<sub>2</sub>, 5 mM EGTA-NaOH, and 0.5 M HEPES-NaOH (pH 6.9) and incubated at 4°C overnight. The fixed tissues were stored in 90% methanol containing 50 mM EGTA at -20°C until use. The fixed and stored tissues were hydrated through decreasing concentrations of methanol and washed with PBS containing 0.2% Tween-20 (PBT). Tissue samples were soaked in PBS containing 2% Tween-20 for two nights, washed with PBT, blocked with PBT containing 5% heat-inactivated goat serum (Wako, Osaka, Japan) and 2% BSA, and incubated with anti-DH[N]<sup>11</sup> at 1:2500 at 4°C overnight. The signal was detected with a Cy3-labeled IgG (Jackson ImmunoResearch Lab, West Grove, PA, USA) diluted to 1:1500 using an FV1000-D confocal microscope (Olympus).

#### **Injection of picrotoxin (PTX) and peptides**

PTX was purchased from Sigma Aldrich (St. Louis, MI, USA). The chemical synthetic Crz peptide was purchased from Abbtotec (Escondido, CA, USA). DH was obtained from Operon Biotechnologies (Tokyo, Japan). PTX and peptides were dissolved in distilled water and peanut oil, respectively, and 10-μL solutions of various doses were injected into pupae a day after pupation through the intersegmental membrane between the second and third abdominal segments.

## cDNA cloning

Eggs were collected the day before hatching. Ovaries were dissected from pupae a day after pupation. Total RNA was extracted from eggs and ovaries using TRI-reagent (Molecular Research Center, Inc., Cincinnati, OH, USA) and then subjected to first-strand DNA synthesis using ReverTra Ace® qPCR RT Master Mix with gDNA Remover (Toyobo, Osaka, Japan). By searching the highly-homologous sequences of each *DHR* and *TRPA1* cDNA for *B. mandarina* in the genomic database SilkBase (<http://silkbases.ab.a.u-tokyo.ac.jp/cgi-bin/index.cgi>), we designed the primers (Supplementary Table S1) to amplify each open reading frame sequence using RT-PCR.

## Electrophysiological assay

TRPA1 channel property was characterized by expressing the channel in *Xenopus laevis* oocytes, and ionic currents were measured by a two-electrode voltage clamp method as previously described<sup>33,34</sup>. In brief, *BmoTRPA1*, *BmaTRPA1a*, and *BmaTRPA1b* were cloned into a pOX plasmid vector, and complementary RNA (cRNA) was synthesized with the mMESSAGE mMACHINE SP6 kit (Thermo Fisher Scientific, Waltham, MA, USA) using a linearized pOX vector containing TRPA1 as a template. Fifty nanoliters of each TRPA1 cRNA (50 or 100 ng/μL) was then injected into defolliculated *Xenopus* oocytes, and ionic currents were observed 2–6 d post-injection. Oocytes were voltage-clamped at –60 mV, and ionic currents were recorded using an OC-725C amplifier (Warner Instruments, Holliston, MA, USA) with a 1-kHz low-pass filter and digitized at 5 kHz using a Digidata 1440 Digitizer (Molecular Devices, San Jose, CA, USA). For thermal stimulation, heated or chilled ND96 bath solutions containing (mM) 96 NaCl, 2 KCl, 1.8 CaCl<sub>2</sub>, 1 MgCl<sub>2</sub>, and 5 HEPES (pH 7.6), were applied by

perfusion. The temperature was monitored with a thermistor located just beside the oocytes using a TC-344B controller (Warner Instruments). Heat-evoked currents for TRPA1 were obtained from *Xenopus* oocytes from two independent preparations, and apparent thermal activation thresholds for TRPA1 were determined with an Arrhenius plot that was generated using Clampfit 10.4 (Molecular Devices) and Origin 9J (OriginLab, Northampton, MA, USA).

### **TALEN construction and screening of KO silkworm**

The TALEN-based mutant lines were constructed according to the previous reports<sup>8,35</sup>. Briefly, TALEN targets were searched using TAL Effector Nucleotide Targeter 2.0 (<https://tale-nt.cac.cornell.edu>) in the coding regions of target genes. DNA constructs containing TAL segments were prepared using a Golden Gate TALEN kit (Addgene, Cambridge, MA, USA). TALEN mRNAs were then synthesized using the mMESSAGE mMACHINE T7 Ultra kit (Ambion, Carlsbad, CA, USA). The mRNA of each TALEN was mixed at a concentration of 0.5 µg/µL for microinjection. Non-diapause eggs of the Kosetsu strain were collected within 1 h after oviposition during the syncytial blastoderm stage. The TALEN mRNA mixture was injected into the eggs using a glass needle (uMPm-02; Daiwa Union, Iida, Japan) attached to a manipulator (kaikopuchu-STDU1; Daiwa Union) and FemtoJet (Eppendorf, Hamburg, Germany).

For screening germline mutagenesis, G<sub>0</sub> adults were mated with *wt.* The oviposited G<sub>1</sub> eggs were collected, and approximately ten eggs from each brood were pooled for genomic DNA extraction using DNAzol® reagent (Thermo Fisher Scientific). The DNA fragment containing the targeted region of interest was amplified by PCR using



Takara Ex Taq (Takara, Tokyo, Japan) and specific primers ([Supplementary Table S1](#)). To test for mutagenesis, the PCR product of *BmoTRPA1* was treated with the Surveyor Mutation Detection kit (IDT, Tokyo, Japan). The mutated PCR products were checked by sequencing. The broods containing mutated sequences were reared, and mutated G<sub>1</sub> adults were crossed with the siblings carrying the same mutation. Homozygous mutants were obtained after confirmation by sequencing of the target region in the G<sub>2</sub> or G<sub>3</sub> egg genome.

#### **DH extraction and titer measurement**

DH was extracted from the hemolymph as previously described<sup>8</sup>. DH extraction was performed at ZT6 (ZT = zeitgeber time, ZT = 0 corresponds to light on). DH levels were measured using a time-resolved fluoroimmunoassay as described previously<sup>8</sup>.

#### **Statistical analysis**

Statistical parameters, including definitions and exact values of *n*, are provided in the relevant figures or corresponding figure legends. Statistical analyses were performed in Microsoft Excel 2011 with the software add-in Toukei-Kaiseki Ver. 3.0 (Esumi, Tokyo, Japan). The significance of differences in diapause egg-inducing activity was evaluated using the Steel-Dwass test. Other data were compared using Student's *t*-tests. Data are expressed as the mean ± SD. *P* < 0.05 was considered significant; ns, non-significant; \*, *P* < 0.05; \*\*, *P* < 0.01; \*\*\*, *P* < 0.001.

#### **Data availability**

The datasets in the current study are available from the corresponding author on

354 reasonable request.

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## 356 References

- 357 1 Nakagaki, M., Takei, R., Nagashima, E. & Yaginuma, T. Cell-Cycles in  
358 Embryos of the Silkworm, Bombyx-Mori - G2-Arrest at Diapause Stage. *Roux Arch Dev*  
359 *Biol* **200**, 223-229 (1991).
- 360 2 Yamashita, O. & Hasegawa, K. in *Comprehensive Insect Physiology,*  
361 *Biochemistry and Pharmacology* Vol. 1 (eds G.A. Kerkut & L.I. Gilbert) 407-434  
362 (Pergamon Press, 1985).
- 363 3 Kogure, M. The influence of light and temperature on certain characters of the  
364 silkworm, Bombyx mori. *Journal of Department of Agriculture, Kyushu University* **4**,  
365 1-93 (1933).
- 366 4 Watanabe, K. Studies on the voltinism of the silkworm, Bombyx mori. *Bull*  
367 *Sericult Exp Sta (Tokyo)* **6**, 411-455 (1924).
- 368 5 Denlinger, D. L., Yocum, G. D. & Rinehart, J. P. in *Insect Endocrinology* (ed L.I.  
369 Gilbert) 430-463 (Academic Press, 2012).
- 370 6 Shimizu, I. & Hasegawa, K. Photoperiodic induction of diapause in the  
371 silkworm, Bombyx mori: location of the photoreceptor using a chemiluminescent paint.  
372 *Physiol Entomol* **13**, 81-88 (1988).
- 373 7 Tsuchida, K. & Yoshitake, N. Effect of different artificial diets on diapause  
374 induction under controlled temperature and photoperiod in the silkworm, Bombyx mori L.  
375 *Physiol Entomol* **8**, 333-338 (1983).
- 376 8 Shiomi, K. *et al.* Disruption of diapause induction by TALEN-based gene  
377 mutagenesis in relation to a unique neuropeptide signaling pathway in Bombyx. *Sci Rep* **5**,  
378 15566 (2015).
- 379 9 Tsuchiya, R. *et al.* Maternal GABAergic and GnRH/corazonin pathway  
380 modulates egg diapause phenotype of the silkworm Bombyx mori. *Proc Natl Acad Sci U*  
381 *SA*, **118**, e2020028118 (2021).
- 382 10 Sato, Y. *et al.* Precursor polyprotein for multiple neuropeptides secreted from  
383 the subesophageal ganglion of the silkworm bombyx-mori - characterization of the  
384 cdna-encoding the diapause hormone precursor and identification of additional peptides.  
385 *P Natl Acad Sci USA* **90**, 3251-3255 (1993).
- 386 11 Hagino, A., Kitagawa, N., Imai, K., Yamashita, O. & Shiomi, K.  
387 Immunoreactive intensity of FXPRL amide neuropeptides in response to environmental  
388 conditions in the silkworm, Bombyx mori. *Cell Tissue Res* **342**, 459-469 (2010).
- 389 12 Yamashita, O., Imai, K., Saito, H., Shiomi, K. & Sato, Y. Phe-X-Pro-Arg-  
390 Leu-NH(2) peptide producing cells in the central nervous system of the silkworm,  
391 Bombyx mori. *J Insect Physiol* **44**, 333-342 (1998).
- 392 13 Homma, T. *et al.* G protein-coupled receptor for diapause hormone, an inducer  
393 of Bombyx embryonic diapause. *Biochem Biophys Res Commun.* **344**, 386-393 (2006).
- 394 14 Sato, A. *et al.* Embryonic thermosensitive TRPA1 determines transgenerational  
395 diapause phenotype of the silkworm, Bombyx mori. *Proc Natl Acad Sci U S A* **111**,  
396 E1249-1255 (2014).
- 397 15 Goldsmith, M. R., Shimada, T. & Abe, H. The genetics and genomics of the

398 silkworm, *Bombyx mori*. *Annu Rev Entomol* **50**, 71-100 (2005).

399 16 Sasaki, C. On the affinity of our wild and domestic silkworms. *Annot. Zool. Jap.*

400 **2**, 33-41 (1898).

401 17 Yoshitake, N. Phylogenetic aspects on the origin of Japanese race of the

402 silkworm, *Bombyx mori* L. *J. Seric. Sci. Jpn.* **37**, 83-87 (1968).

403 18 Yukuhiro, K., Sezutsu, H., Itoh, M., Shimizu, K. & Banno, Y. Significant levels

404 of sequence divergence and gene Rearrangements have occurred between the

405 mitochondrial Genomes of the wild mulberry silkmoth, *Bombyx mandarina*, and its close

406 relative, the domesticated silkmoth, *Bombyx mori*. *Mol Biol Evol* **19**, 1385-1389 (2002).

407 19 Miyata, T. A generic revision of the Japanese Bombycidae, with description of a

408 new genus (Lepidoptera). *Tinea* **8**, 190-199 (1970).

409 20 Murakami, A. & Imai, H. T. Cytological evidence for holocentric chromosomes

410 of the silkworms, *Bombyx mori* and *B. mandarina*, (Bombycidae, Lepidoptera).

411 *Chromosoma* **47**, 167-178 (1974).

412 21 Yokoyama, T. Reproduction and natural enemies of the wild mulberry silkmoth

413 (*Bombyx mandarina*). *Sanshi-Konchu Biotec* **88**, 25-38 (2019).

414 22 Xia, Q. Y. *et al* Complete resequencing of 40

415 genomes reveals domestication events and genes in silkworm (*Bombyx*). *Science* **326**,

416 433-436 (2009).

417 23 Umeya, Y. Embryonic hibernation and diapause in insects from the viewpoint of

418 the hibernating-eggs of the silkworm. *Bull. Sericul. Exp. Sta.* **12**, 393-480 (1946).

419 24 Yamashita, O. & Yaginuma, T. in *Insects at Low Temperature* (eds Jr. R. E. Lee

420 & D. L. Denlinger) 424-445 (Chapman and Hall, 1991).

421 25 Komoto, N. & Tomita, S. Distribution and seasonal occurrence of the wild

422 mulberry silkmoth (*Bombyx mandarina*) in Japan. *Sanshi-Konchu Biotec* **88**, 7-23

423 (2019).

424 26 Ohmura, S. Research on the behavior and ecological characteristics of the wild

425 silkworm, *Bombyx mandarina*. *Bull. Seric. Exp. Sta. Jpn.* **13**, 79-130 (1950).

426 27 Kobayashi, J. Effects of photoperiod on the induction of egg diapause of

427 tropical races of the domestic silkworm, *Bombyx-Mori*, and the wild silkworm,

428 *Bombyx-mandarina*. *JARQ Jpn Agric Res Q* **23**, 202-205 (1990).

429 28 Xu, W. H., Sato, Y. & Yamashita, O. Molecular characterization of the cDNA

430 encoding diapause hormone and pheromone biosynthesis activating neuropeptide in

431 *Bombyx mandarina*. *J. Seric. Sci. Jpn.* **68**, 373-379 (1999).

432 29 Li, T. B. *et al*. Diverse sensitivities of TRPA1 from different mosquito species to

433 thermal and chemical stimuli. *Sci Rep-Uk* **9**; ARTN 2020010.1038/s41598-019-56639-w

434 (2019).

435 30 Zhong, L. X. *et al*. Thermosensory and Nonthermosensory Isoforms of

436 *Drosophila melanogaster* TRPA1 Reveal Heat-Sensor Domains of a ThermoTRP

437 Channel. *Cell Rep* **1**, 43-55 (2012).

438 31 Denlinger, D. L., Hahn, D. A., Merlin, C., Holzapfel, C. M. & Bradshaw, W.

439 E. Keeping time without a spine: what can the insect clock teach us about seasonal

440 adaptation? *Philos Trans R Soc Lond B* **372**; ARTN 2016025710.1098/rstb.2016.0257

441 (2017).

442 32 Kobayashi, J. Effects of photoperiod on the duration of pupal stage of the wild

443 silkworm, *Bombyx mandarina* Moore. *Wild Silkmooths '89-'90*, 57-64 (1991).

444 33 Saito, S. *et al* Heat and noxious chemical sensor, chicken TRPA1, as a target of

445 bird repellents and identification of its structural determinants by  
 446 multispecies functional comparison. *Mol Biol Evol* **31**, 708-722(2014).  
 447 34 Saito, S., Saito, C. T., Nozawa, M. & Tominaga, M. Elucidating the functional  
 448 evolution of heat sensors among *Xenopus* species adapted to different thermal niches by  
 449 ancestral sequence reconstruction. *Mol Ecol* **28**, 3561-3571 (2019).  
 450 35 Takasu, Y., Tamura, T., Sajwan, S., Kobayashi, I. & Zurovec, M. The use of  
 451 TALENs for nonhomologous end joining mutagenesis in silkworm and fruitfly. *Methods*  
 452 **69**, 46-57 (2014).  
 453  
 454

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**Author contributions**

K.S. designed the research. K.Y., S.S., M.S., M.K., Y.T., H.S., Y.K., M.T., A.K., and K.S. performed the research. K.Y., S.S., and K.S. analyzed the data. T.Y., S.S., and K.S. drafted the manuscript with support from all authors.

**Competing interests**

The authors declare no competing financial interests.

## Figure legends

**Figure 1.** Temperature and photoperiodic diapause induction in *Bombyx mori* and *B. mandarina*. **(a)** The typical larvae and eggs of the wild mulberry silkworm *B. mandarina*. Eggs were incubated at 25DD; after that, larvae were reared at 16L8D or 8L16D. Progeny eggs hatched on 10 d after oviposition under the 16D8D condition, whereas progeny eggs never hatched by 30 d after oviposition under the 8L16D condition. **(b)** Effects of temperature and photoperiodic conditions during embryonic and larval stages on diapause egg-inducing activity in the Kosetsu strain of *B. mori* and a Fuchu population of *B. mandarina*. After eggs were incubated under the 25DD or 15DD condition, larvae were reared on 24L0, 16L8D, and 8L16D conditions. Each bar represents the mean  $\pm$  SD of 5–20 animals. Significant differences represented in *B. mori* versus *B. mandarina* are in the same condition. ns, non-significant; \*\*,  $P < 0.01$ ; \*\*\*,  $P < 0.001$ . **(c)** Thionin staining of embryos 30 d after oviposition of domestic silkworm *B. mori*, and 30 d after oviposition of *B. mandarina* in both 8L16D and DH-injected silkworm in 16L8D. Scale bar = 200  $\mu$ m.

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**Figure 3.** Cloning and functional analysis of *Bombyx mori* and *B. mandarina* TRPA1. (a) Schematic representations of the *B. mori* TRPA1 (*BmoTRPA1*), *B. mandarina* TRPA1a (*BmaTRPA1a*), and TRPA1b (*BmaTRPA1b*). Boxes and lines represent exons and introns, respectively. Black, blue, orange, gray, and red boxes represent domains of untranslated regions, ankyrin repeat, linker region, transmembrane (1~6), and pore-loop, respectively. The red triangle indicates the TALEN target site. (b) Sequence alignment of the linker region between TRPA1s. (c–e) Arrhenius plots for the heat-induced *BmoTRPA1*, *BmaTRPA1a*, and *BmaTRPA1b* current shows a clear flex point in temperature dependency. Each temperature threshold for TRPA1 activation was defined as when the two linear-fitted lines crossed (i.e., a flex point). (f) Comparison of threshold temperature for each TRPA1. Each bar represents the mean  $\pm$  SEM of 8–11 samples. Significant differences versus *BmoTRPA1* were evaluated. ns, non-significant.

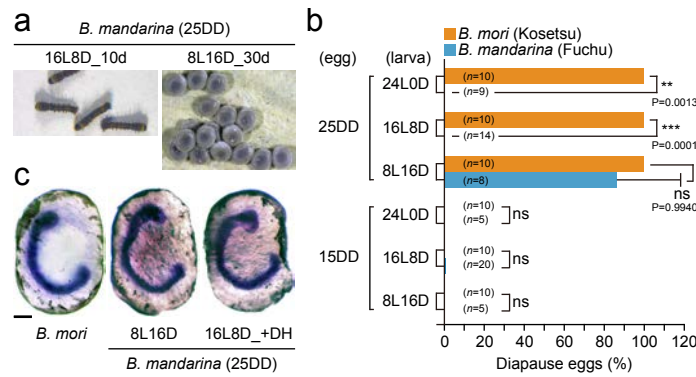
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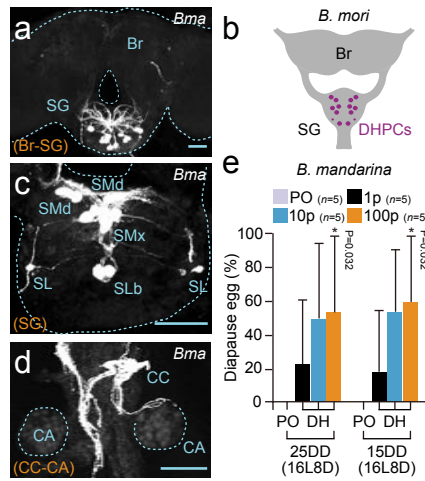
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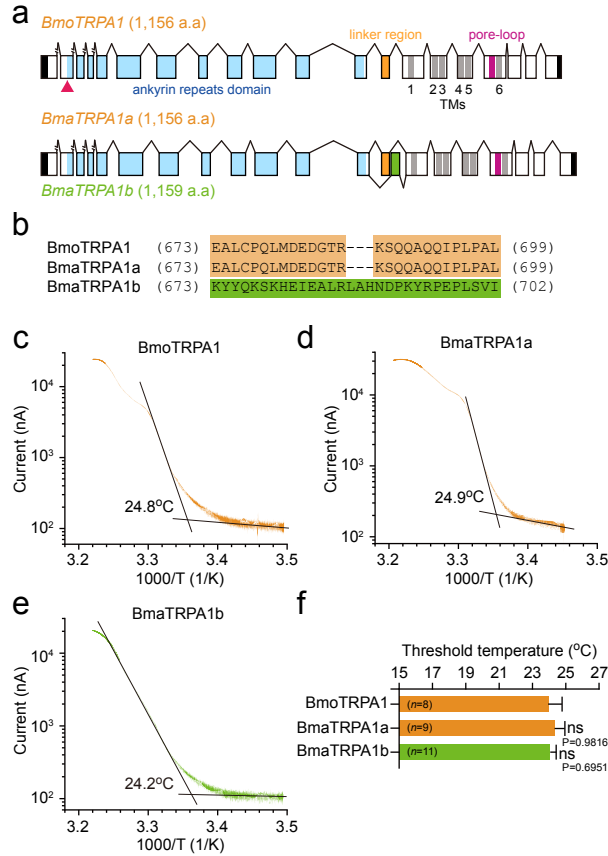




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## Supplementary Table S1

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### List of primers\*

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#### a. For cDNA cloning

- |          |  |
|----------|--|
| 1. DHR   | F: 5'-CGTCACGTAACGTCAACATTGCGGTACGC-3'<br>R: 5'-TAATTACTTACTAAACGGAATGTATTTG-3'  |
| 2. TRPA1 | F: 5'-GTCATGACATTTGTGGAACAACGGCCTA-3'<br>R: 5'-TCAAGTCGACTTGTTAAATGATGCAGCTCG-3' |
- 

#### b. For screening of the $\Delta TRPA1$ mutant

- |  |
|--|
| F: 5'-AGTTTTTTGTACAGGCTGCTGAAAGCGGA-3' |
| R: 5'-TCTCGGTTTGCATCGCCAATAAGC-3'      |
- 

\*F; Forward primer, R; Reverse primer.

**a**

```
BmoDHR; TDMKDESDRGASRGALWFGPRL 24
BmaDHR; TDMKDESDRGASRGALWFGPRL 24
*****
```

**b**

```

                                     TM1
BmoDHR; MNS--ETINDTANASRPVDSTRVFGPQRTDLYIVLPITIIYTFIFVSGLLGNIFTCIVIV 58
BmaDHR; MNSLNETLNDTANASRPVDSTRVFGPQRTDLYIVLPITIIYTFIFISGLLGNIFTCIVIV 60
*** **

          TM2                      TM3
BmoDHR; RNKNLHTATNYLFLSLAISDLLLVSGMPQEMYSIWSKWPYVFGHTFCVIRGLAAETSTN 118
BmaDHR; RNKNLHTATNYLFLSLAISDLLLVSGMPQEMYSIWSKWPYVFGHTFCVIRGLAAETSTN 120
*****

                                     TM4
BmoDHR; ASVLTITLFTIERYLAICHFFVSHKMSKLSRAVKHVLLWVAALALALPQALQFGIRQYQ 178
BmaDHR; ASVLTITLFTIERYLAICHFFVSHKMSKLSRAVKHVLLWVAALALALPQALQFGIRQYQ 180
*****

          TM5
BmoDHR; GVIMCLQTRVIEHSFEISTFLFPLAPMLITVLYSFIGLKLREKSNVKEQNQNDFESSI 238
BmaDHR; GVIMCLQTRVIEHSFEISTFVFFLAPMLITVLYSFIGLKLREKSNVKEQNQNDFESSI 240
*****

          TM6
BmoDHR; RYSHKMCRKPSQSTRRVIKMLVAVVVAFFICWAPFHAQRLVAIYGTNENHLAKSPILFSV 298
BmaDHR; RYSHKMCRKPSQSTRRVIKMLVAVVVAFFVCWAPFHAQRLVAIYGTNENHLAKSPILFSV 300
*****

          TM7
BmoDHR; YLFLTYISGIFYYMSTCINPILYHIMSNKFRDAFKMTLCCCGTRNDTAVKRSSYTAMAFV 358
BmaDHR; YLFLTYISGICYMSTCINPILYHIMSNKFRDAFKMTLCSGTRNDTAVKRSSYTAMAFV 360
*****

BmoDHR; RHPTSSGTSNSGNSIRNETNLQSKTRRTNGRDKILNDAHVCNGTLSSAAVGKPDSTRNG 418
BmaDHR; RNPTSSGTSNSGNSIRNETNLQSKTRRTNSRDKILNDAHVCNGTLSSAAVGKPDSTRNG 420
*

BmoDHR; DRPLDRNLINETYFNTNC 436
BmaDHR; DRPLDRNLINETYFNTNC 438
*****

```

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# Figures

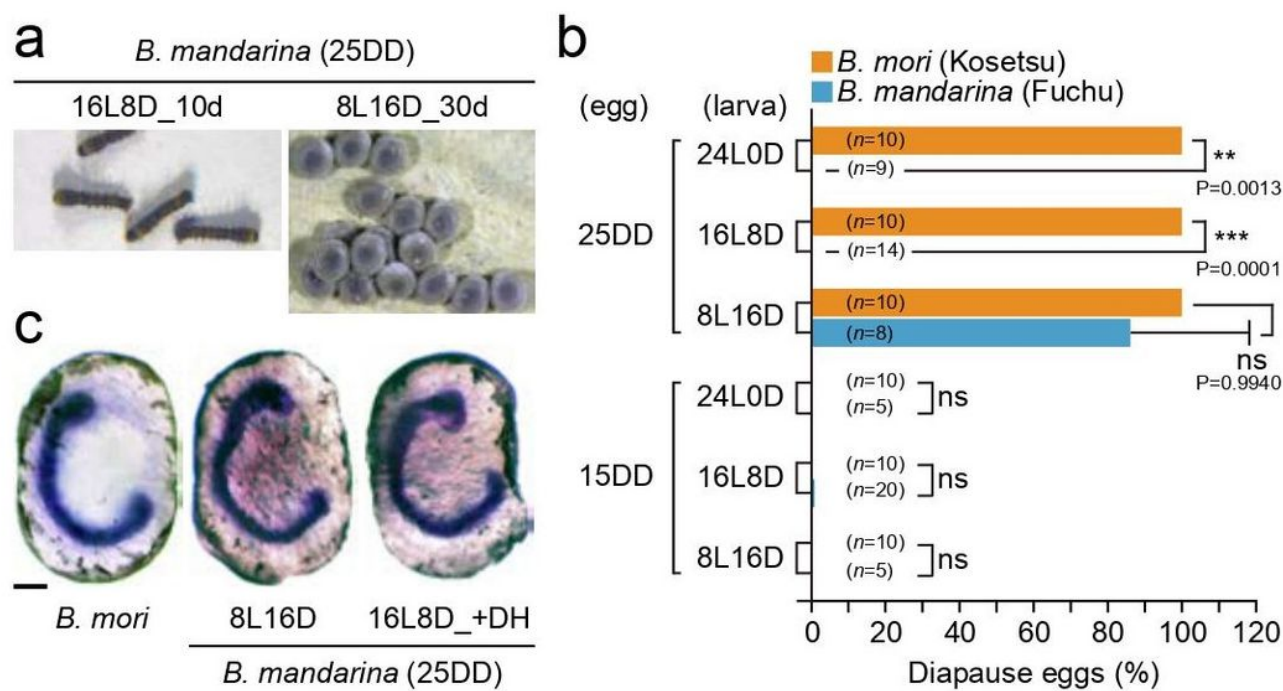
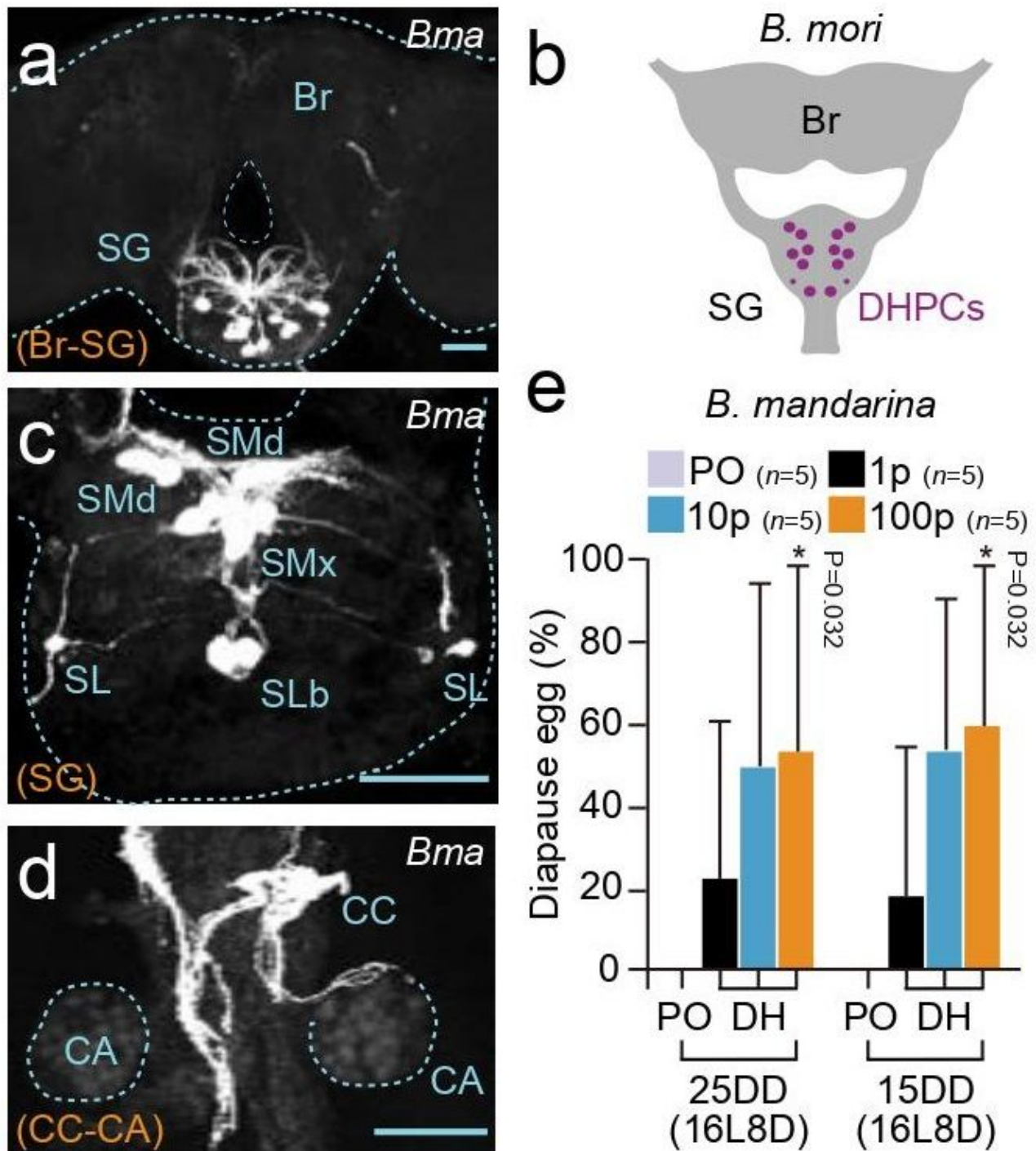


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Diapause egg-inducing activity of DH injection. Peanut oil (PO) or each DH peptide concentration (1, 10, or 100 pmol) was injected into *B. mandarina* pupa reared under the 16L8D condition from 25DD and 15DD embryos, and diapause egg-inducing activity was measured. Each bar represents the mean  $\pm$  SD of five animals. Asterisks indicate significant differences versus PO injection. \*,  $P < 0.05$ .

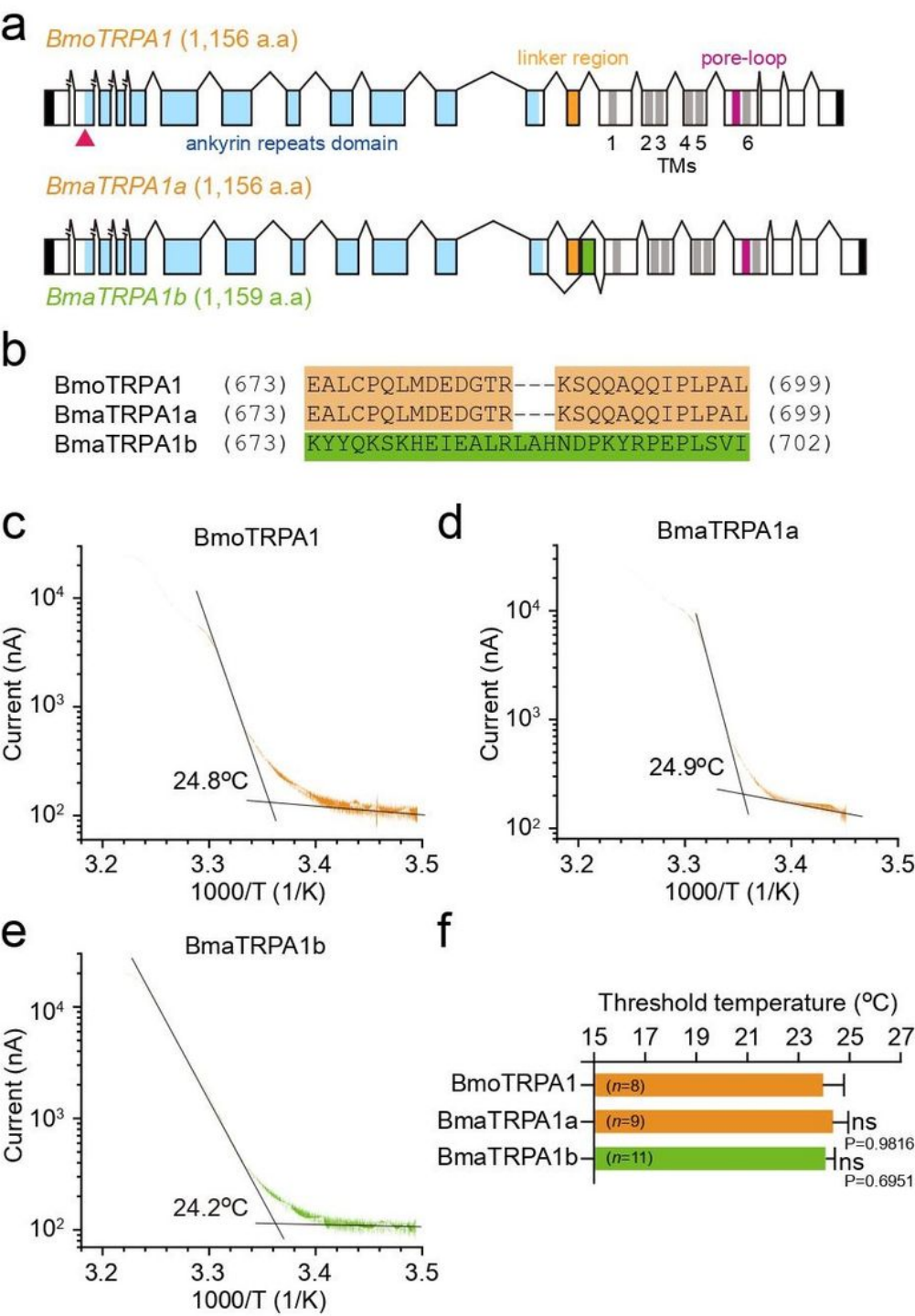


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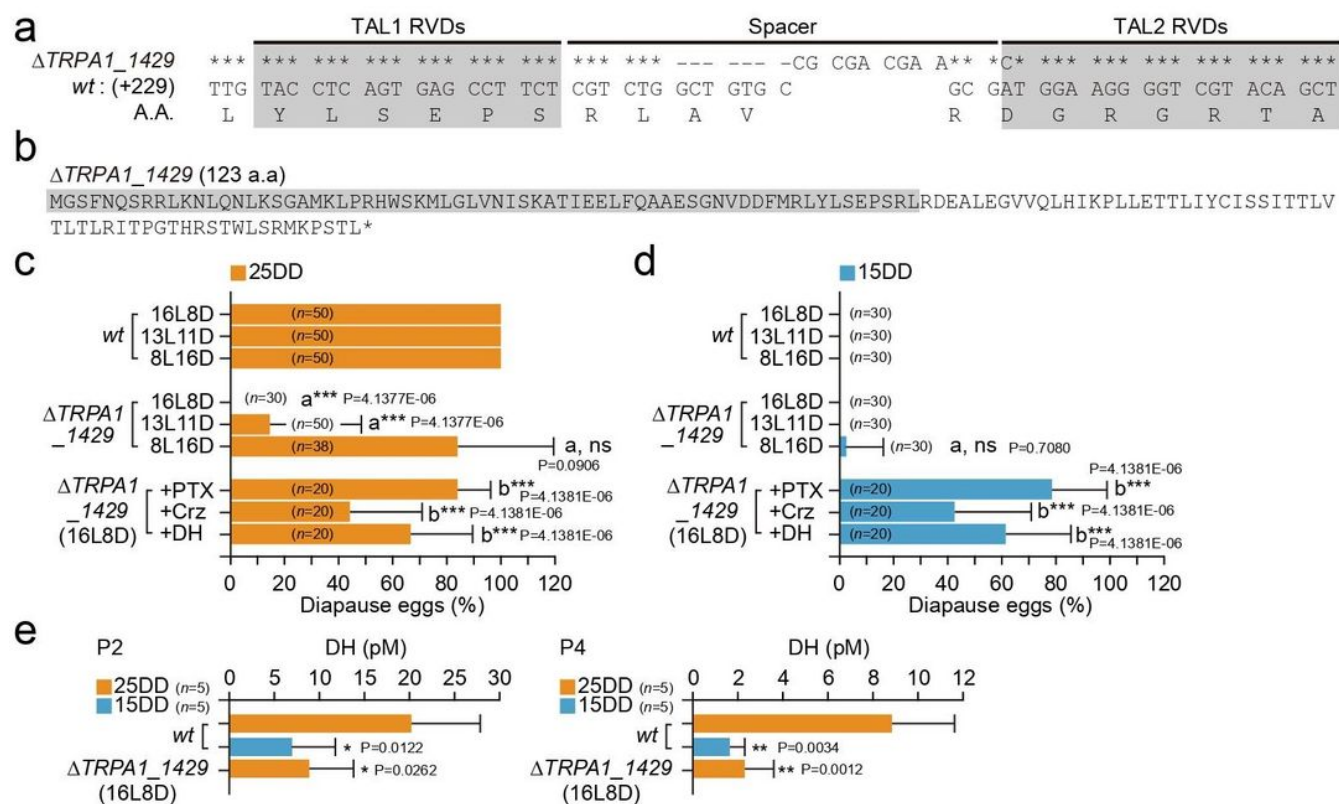


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## Supplementary Files

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