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A Bombyx homolog of ovo is a segmentation gene that acts downstream of Bm-wnt1(Bombyx wnt1 homolog)

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Keywords

Bombyx, ovo, wnt1, RNAi, segmentation, pair-rule genes

Abstract

Insect embryogenesis is divided into long and short/intermediate germ types. The long germ type may exhibit Drosophila-like hierarchical segmentation mechanisms, whereas the short/intermediate type assumes some repeating mechanisms that are considered to be ancestral. Embryogenesis in Bombyx mori possesses both characteristics. Here, Bombyx ovo homolog (Bm-ovo) was identified as a gene involved in segmentation. ovo is a Drosophila gene that encodes a zinc finger transcription factor and studies on its homolog functions in other systems have suggested that it acts as a switch to enable the initiation of differentiation from a progenitor cell state. This is the first description for ovo homologs being involved in insect segmentation. Bm-ovo is expressed dynamically during embryogenesis in a pattern that resembles that of gap and pair-rule genes. In Bm-ovo RNAi knockdown embryos, posterior segmentation does not proceed. In addition, defects in anterior segments are observed. In Bm-wntl knockdown embryos, the Bm-ovo expression pattern was changed, suggesting that Bm-wntl is an upstream regulator of Bm-ovo. The involvement of Bm-ovo may represent a novel ancestral step under the control of wnt genes in insect segmentation: this step may resemble those operating in cell differentiation processes.

Introduction

In insect embryos, maternal, gap, and pair-rule genes are successively expressed to produce segments along the anterior-posterior (AP) axis. Maternal genes dictate the AP axis. Gap genes subdivide embryos into broad domains according to their expression profiles. The expression of seven to eight stripes of pair-rule genes serves as a template for future segments. Insect embryogenesis has been largely categorized into long and intermediate/short germ types (Davis and Patel, 2002; Liu and Kaufman, 2005; Peel et al., 2005). In the long germ type, as represented by *Drosophila*, the combinatorial input of maternal and gap genes directs the expression of the individual stripes of pair-rule genes, which leads to the nearly simultaneous expression of pair-rule stripes, i.e., segment formation (Pankratz and Jäckle, 1993). In contrast, in the short/intermediate type, posterior segments are sequentially added, in which some repeating segmentation mechanisms are likely to operate. In the short germ-type insect Tribolium, the wave-like expression of pair-rule genes even-skipped (eve) and odd-skipped (odd) has been observed in the pre-segmental area (Sarrazin et al., 2012; El-Sherif et al., 2012). The pair-rule genes, eve, odd, and runt (run) interact positively and negatively to form a circuit that appears to be responsible for the striped expression of these genes. Thus, the wave-like expression of a subset of pair-rule genes may be passed onto the pair-rule gene circuit, leading to the production of segments (pair-rule gene stripes) in a clock-like manner (Choe et al., 2006; Lynch et al., 2012).

It is commonly acknowledged that the latter *Tribolium*-like segmentation mode is ancestral, while the former *Drosophila*-like hierarchical segmentation mode is derived. Recent evidence suggests that both modes may operate in some species (Rosenberg et al., 2014), such as *Bombyx mori*.

B. mori is a lepidopteran insect, the segmentation mechanisms of which have remained elusive. Its embryogenesis has long and short/intermediate-like features. The egg has a large embryonic primordium, within which individual segments are fate mapped without a growth zone early in embryogenesis similar to long germ insects (Myohara, 1994). However, when pair-rule gene interactions were examined, *B. mori* exhibited *Drosophila-* and *Tribolium*-like characteristics (Nakao, 2015). Furthermore, the roles of *Bombyx wnt1/wingless* and the *Krüppel* homolog resemble those in short/intermediate insects: embryos show truncated phenotypes in the posterior region when these gene

(family) activities are perturbed (Beermann et al., 2011; Bolognesi et al., 2008, 2009; Miyawaki et al., 2004; Nakao, 2010, 2015; Yamaguchi et al., 2011; Cerny et al., 2005; Liu and Kaufman, 2004; Mito et al., 2006). Embryonic Bm-wnt1 is considered to have two functions: its organizing function in posterior development, as reflected in the knockdown phenotype as described above, and its function as a segment polarity gene, similar to that in *Drosophila*. During normal embryogenesis, *Bm-wnt1* is expressed in the large posterior domain in the early stages, the expression domain then recedes posteriorly, and segmental stripes begin to appear in an anterior to posterior sequence (Nakao, 2010). Posterior expression appears to be responsible for posterior organization functions, while segmental expression appears to exhibit a segment polarity function that has not yet been experimentally validated in Bombyx. Furthermore, the addition of supernumerary posterior segments is induced at the posterior terminus in a manner that resembles that of short/intermediate insects by the knockdown of the hunchback homolog (Bm-hb) (Nakao, 2016). These findings strongly suggest that the short/intermediate type mode of segmentation is operating in normal Bombyx embryogenesis.

An ovo homolog from B. mori (Bm-ovo) was described herein. ovo is a Drosophila gene that encodes an evolutionarily conserved zinc finger transcription factor with various biological functions. It is required in the female germ line for proper oogenesis: the mutants display egg chambers filled with excess undifferentiated germ cells, an ovarian tumor phenotype (Oliver, 1987). ovo is also involved in early germ line development: the transcript is deposited in the germ plasm as a maternal factor, regulating the expression of vasa (Yatsu et al., 2008). In the zygote, often referred to as shavenbaby (svb) at this stage, it is necessary and sufficient to cell-autonomously direct the formation of denticles (trichome-cytoplasmic extrusion of epidermal cells) in the ventral epidermis of the embryo abdomen, in which there is a segmentally repeating denticle pattern separated by naked cuticle, thereby contributing to the binary choice to produce either a naked cuticle or denticles. The upstream regulation of svb has been investigated. The high mobility group (HMG)-domain protein SoxNeuro (SoxN) is necessary and sufficient to cell-autonomously direct the expression of svb. SoxN, in turn, receives positive and negative inputs from the epidermal growth factor receptor (Egfr) ligand Spitz (Spi) and Wingless (Wg), respectively. These mechanisms result in the restriction of trichome-producing cells. The closely related protein of Spi, Dichaete is co-regulated

with Spi and has a partially redundant function in the activation of *svb*, albeit to a lesser extent. However, these regulatory relationships are not strictly hierarchical, but complex feedback mechanisms are involved (Overton et al., 2007).

Apart from *Drosophila*, *ovo* homologs are isolated from diverse animals. The mouse homolog also controls germline and epidermis differentiation. In *C. elegans*, the *ovo* homolog *lin-48* is required for the development of hindgut. These findings point to the role of *ovo* homologs in the differentiation and maintenance of specific cell types (Wieschaus, 1984; Oliver, 1987; Dai, 1998; Johnson, 2001). More recently, human homolog of *ovo* (*OVOL1*) was shown to regulate the transition of progenitor to differentiated trophoblast cells (Renaud et al., 2015).

In the present study, the embryonic functions of *Bm-ovo* were examined. *Bm-ovo* expression during the embryonic stage resembled that of gap and pair-rule genes. Embryonic *Bm-ovo* RNAi embryos showed segmentation defects in the gnathal/thoracic region and posterior abdomen. In the posterior body part, the abnormal expression of some pair-rule genes that are crucial for establishing pair-rule gene expression patterns was observed: the posterior part did not produce pair-rule stripes, but a broad band of expression was observed in the affected region for the pair-rule genes examined. In *Bm-wnt1* knockdown embryos, alternation in *Bm-ovo* expression pattern was observed, suggesting that *Bm-wnt1* is an upstream regulator of *Bm-ovo*. This interaction may be related to those operating in *Drosophila* embryonic epidermal patterning as described above.

The involvement of *Bm-ovo* may represent a novel ancestral step under the control of *wnt* genes in insect segmentation, and this step may resemble those operating in cell differentiation processes.

Materials and Methods

Silkworm strains, rearing, and development

The *B. mori* strain *pnd-2* was used in this study. Silkworms were reared on an artificial diet (Nippon Nosanko). Refer to Nagy et al. (1994) for a general description of early *Bombyx* development.

Identification and isolation of Bm-ovo cDNA

The Drosophila ovo sequence was used to search the Bombyx cDNA database (Mita et al., 2003) for Bombyx ovo homologs and one wing-disk-derived cDNA was identified. cDNA was obtained and initial analyses were performed using this cDNA. The annotated genome database (KAIKObase; Mita et al., 2004) subsequently became available, which revealed that the open reading frame of *Bm-ovo* comprises four exons (E1, E2, E3 and E4; gene ID: BMgn000987). The cDNA clone described above was found to lack the second exon. To obtain cDNA species expressed during early embryogenesis, a PCR amplification procedure was employed. Total RNAs in embryos 14 and 20 hours AEL were prepared using TRIZOL reagent (Invitrogen). Single-strand cDNAs were synthesized using total RNAs as templates by PrimeScriptTMII 1st strand cDNA Synthesis Kit (TAKARA) according to the manufacturer's instructions. With these cDNAs, PCR amplification was performed using PrimeSTAR GXL DNA Polymerase (TAKARA). A primer pair was set at the sequence near the start and stop codons, respectively, so as to amplify the full open reading frame. PCR conditions were 40 cycles at 98°C for 10 s; at 55°C for 15 s; at 68°C for 3 min. The primer pair was the *Bm-ovoCp*, 5'-primer and *Bm-ovoE3-4p*, 3'-primer (see below). The amplified product was analyzed using 1% agarose gel electrophoresis. The results obtained revealed the predominant amplification of an approximately 2.6-kbp product from both templates (Supplementary Fig. 1). The amplified product was cloned and sequenced. The sequence analysis revealed that it had all four exons. Subsequent analyses were performed using this cDNA.

In situ hybridization

Fixation and *in situ* hybridization were performed as previously described (Nakao, 1999, 2010). Probes for *Bm-eve* and *Bm-wnt1* were previously described (Nakao, 2010). Probes for *Bm-ovo*, designated as *Bm-ovoCp*, *Bm-ovoE1p*, *Bm-ovoE2p* and *Bm-ovoE3-4p*, were synthesized using cloned cDNAs. cDNAs were PCR fragments amplified using the primers described below. The primer pairs used for the amplification of these fragments were as follows: *Bm-ovoCp*, 5'-primer: 5'-GGGGGATCCAGTCCTAACGAAGCGGCCAA-3', 3'-primer: 5'-

CCCAAGCTTTTTATACGGTCTGACTCCGG-3'; Bm-ovoElp, 5'-5'-primer: 3'-primer 5'-GGGGGATCCAGTCCTAACGAAGCGGCCAA -3', CCCAAGCTTCAGTGCATTCCTTTTCTTTATCC -3'; Bm-ovoE2p, 5'-primer: 5'-5'-GGGGGATCCCGCACAAAAGAAACTAGACG -3', 3'-primer CCCAAGCTTAGCACTAAAACAGGTCGTGC -3'; Bm-ovoE3-4p, 5'-primer: 5'-GGGGGATCCTAGGACTACCAGCAGAGCTT -3', 3'-primer 5'-CCCAAGCTTAATTGTGTACTGGCATGGGC -3'. The Bm-ovoCp primer pair was used for amplification from the cloned cDNA template (fufe-P20 F P18), and the amplified product comprised approximately 0.3-kb exon 1 and 0.3-kb exon 3 (see the Results section). Bm-ovoE1p, E2p and E3-4p primer pairs were used for amplification from the cDNA prepared using PrimeScriptTM II 1st strand cDNA Synthesis Kit (TAKARA) and total RNA at 20 hours AEL. The amplified products were approximately 0.3, 0.5, and 0.6 kbp, which corresponded to parts of exons 1, 2, and 3-4, respectively. After amplification, cDNAs were cloned into pBluescript vectors. In order to obtain probes, plasmid DNA was cut with an appropriate enzyme and RNA probes were synthesized using either T3 or T7 polymerase, depending on the direction of the insert using DIG RNA Labeling Kit (SP6/T7) (Roche).

RNAi

The RNAi procedure was described previously (Nakao, 2012). The templates used for in vitro transcription were PCR fragments of the corresponding genes, flanked by T7 promoter sequences. Regarding Bm-ovo RNAi, three different dsRNAs, designated as Bm-ovoCRNAi, Bm-ovoE1RNAi and Bm-ovoE3-4RNAi, were used in analyses. The amplification primer pairs used for were as follows: Bm-ovoCRNAi, 5'-TAATACGACTCACTATAGGGAGAAGTCCTAACGAAGCGGCCAA-3', 5'-TAATACGACTCACTATAGGGAGATTTATACGGTCTGACTCCGG-3'; 5'-Bm-ovoE1RNAi, TAATACGACTCACTATAGGGAGAAGTCCTAACGAAGCGGCCAA-3', 5'-TAATACGACTCACTATAGGGAGACAGTGCATTCCTTTTCTTTATCC-3'; Bm-ovoE3-4RNAi, 5'-TAATACGACTCACTATAGGGAGATAGGACTACCAGCAGAGCTT-3', 5'-TAATACGACTCACTATAGGGAGAAATTGTGTACTGGCATGGGC. Bm-ovoC. ovoE1, and E3-4RNAi targets corresponded to the regions amplified by the Bm-ovoCp, Bm-ovoE1p, and Bm-E3-4p primer pairs described above, respectively. For Bm-wnt1 RNAi, two independent dsRNA corresponding to different part of the gene was used for the The follows: Bmwnt1RNAi1. analysis. primer pairs were as 5'-TAATACGACTCACTATAGGGAGAGCAGAATGAAGTGTCTGTGG-3', 5'-TAATACGACTCACTATAGGGAGAAAGCCGATGTTGTCGCTGCA-3', Bmwnt1RNAi2, 5'-TAATACGACTCACTATAGGGAGAAGGGAATTCGTTGATACCGG-3', 5'-TAATACGACTCACTATAGGGAGAACCTCGCAACACCAATGGAA-3'.

Results

Identification of a Bombyx ovo homolog

A recent study described the *ovo* homolog from *Bombyx* (*Bm-ovo*); multiple forms of spliced variants were differentially expressed among the tissues examined (Xue et al., 2014). The identification of *Bm-ovo* in the present study was performed independently in our laboratory. After cDNA database search (Mita et al., 2003), a cDNA derived from wing-disk-expressed mRNA was obtained (fufe-P20_F_P18). Initial analyses were performed using a probe (*Bm-ovoCp*) or double-stranded RNA (*Bm-ovoCRNAi*) made from this cDNA as a template. However, a subsequent analysis revealed that this mRNA species lacked exon E2 and was poorly represented in early embryonic stages (see **Materials and Methods**). After obtaining embryonically expressed cDNA, which comprised exons E1, E2, E3 and E4, analyses were also performed using a few mutually independent probes or dsRNA in addition to the analyses described above. The results of these analyses were essentially the same.

Bm-ovo embryonic expression

A gene expression analysis of *Bm-ovo* was performed by *in situ* hybridization using the probes, *Bm-ovoCp*, *E1p*, *E2p* and *E3-4p* (see **Materials and Methods**). Irrespective of probes used, the results were essentially the same (Supplementary Fig. 2). Just after egg laying, weak expression was observed in the germ anlage, suggesting its maternal load

(Fig. 1A, B). Regional heterogeneities in expression were not detected within the germ anlage, suggesting that the transcript did not accumulate in the putative germ plasm, although the existence of germ plasm was suggested by *Bm-nanosO* (*Bombyx nanosO*) expression pattern (Nakao et al., 2008). Specific accumulation in the germ line was not observed during the time period examined in the present study, i.e., up to 5 days AEL (After Egg Laying), which was in contrast with previous findings obtained using Drosophila (Yatsu et al., 2008). Expression was dynamic more than 22 hours AEL and resembled that of the gap and pair-rule genes. At 22 hours, expression was detected in one anterior domain (head) and one large posterior domain (Fig. 1C). Later, the posterior domain resolved into three strong stripes (bands) and two weak stripes (bracketed) (Fig. 1D, E). The three strongly expressed stripes remained thereafter, and overlapped with Bm-eve stripes #2, #6, and #8 (Fig. 1F, G). Although it could not be determined due to technical difficulties which pair-rule stripes the weak Bm-ovo stripes corresponded to or which pair-rule stripes the broad strong Bm-ovo stripes exactly cover, a (morphological) comparison of single- and double-stained embryos suggested that stripes #1, #3, and #7 escaped Bm-ovo coverage at the later stages. However, the expression pattern of Bm-ovo suggested that the Bm-ovo posterior expression domain covered all the Bm-ovo stripes at least once, except for stripe #1. As the development proceeded, the background signal intensity transiently increased, region-specific staining became obscured (Fig. 1H), and background staining then decreased again (Fig. 1I).

Analysis of Bm-ovo functions using embryonic RNAi

Embryonic RNAi was used to analyze *Bm-ovo* functions. dsRNAs were injected into embryos at $0\sim2$ hours AEL. As described above, three dsRNAs were used to examine phenotypes. Target gene expression was markedly reduced for all the dsRNAs used, suggesting that the RNAi procedure was effective (Supplementary Fig. 3).

The phenotypes of knockdown embryos were examined for their morphologies at 72 hour AEL. In RNAi-treated embryos, anomalies were observed in the anterior and posterior of embryos (Fig. 2). Anterior phenotypes represented defects in the morphology of the region encompassing gnathal/thoracic segments, with the mesothoracic segment being the most frequently affected. In these regions, the observed

phenotypes included irregular morphologies or the deletion of segments and/or appendages, or fused or disorganized gnathal/thoracic segments (Fig. 2B, C). In the posterior, the fusion of posterior abdominal segments was observed for all dsRNA species: in the most severe cases, a total of six abdominal segments were observed (Fig. 2B, C, Table 1). The identities of the remaining abdominal segments were not entirely clear. However, based on the observation that three proleg-bearing segments remained in RNAi treated embryos (Fig. 2B), the identities of A1 to A5 appeared to be preserved in the correct order (in the wild-type, prolegs were observed in A3-A6, also see below). In both phenotype classes, the severity of the phenotype appeared to differ between the dsRNAs used (Table 1); for example, the number of abdominal segments was significantly lower in *Bm-ovoE1RNAi* or *Bm-ovoE3RNAi* embryos than in *Bm-ovoCRNAi* embryos (P<0.05), reflecting the incomplete nature of the RNAi knockdown. Therefore, the knockout phenotype may differ from those observed here by the RNAi knockdown.

In order to elucidate the cause of these morphological consequences, embryos were analyzed molecularly. Since pair-rule genes were responsible for segment formation, the expression of three key pair-rule genes, *Bombyx even-skipped (Bm-eve)*, *runt (Bm-run)*, and *odd-skipped (Bm-odd)* was examined at 23~28 hours AEL when the full complement of each pair-rule gene stripe was expressed in wild-type embryos (Fig. 3A, C, E; Nakao, 2015). The most notable change was the broad expression of these pair-rule genes in the posterior region without the resolution of stripes after stripe #5, which is consistent with posterior abdominal segment fusion morphologies (*Bm-eve*, n=29/31; *Bm-run*, n=27/28; *Bm-odd*, n=50/54). In addition, the weak expression of the *Bm-eve* stripe #3 (and less frequently, stripe #2) was sometimes observed (n=15/21), together with the weak expression of *Bm-run* and *Bm-odd* stripe #2 (*Bm-run*, n=5/25; *Bm-odd*, n=16/51; Fig. 3B, D, F, Supplementary Fig. 4). However, it is important to note that although the intensity of expression was changed, patterning in the anterior region was essentially not affected by *Bm-ovo* RNAi, which is in contrast to posterior patterning in the affected embryos.

Bm-ovo is regulated by Bm-wnt1

The present findings show that Bm-ovo is involved in Bombyx body segmentation.

Bm-wnt1 also functions in the establishment of segments. These homologs are known to interact in Drosophila embryonic epidermal patterning as described in the Introduction section. Therefore, embryonic RNAi was used to test whether Bm-ovo and Bm-wntl are interact and, if so, how. Bm-ovo RNAi was performed as described above and Bm-wntl expression in the resulting embryos was compared with that in wild type embryos. As examined at ~23h AEL, Bm-wnt1 expression in RNAi-treated embryos was similar to that in wild type embryos (n=14/14; Fig. 4A, B). However, at approximately 26h AEL, when more anterior segmental stripes of Bm-wntl became visible, these stripe boundaries in treated embryos were more irregular, except for a few anterior-most stripes (n=20/20; black arrows in Fig. 4D), than those in wild type embryos, whereas differences were not observed in the posterior expression domain (n=34/34; white arrows in Fig. 4A-D). Next, Bm-wntl RNAi was performed to examine changes in Bm-ovo expression. The in situ hybridization of RNAi treated embryos using the Bm-wntl probe indicated that the target gene expression was markedly reduced, suggesting that the RNAi was effective (Supplementary Fig. 3). A previously study reported that Bm-wntl knockdown resulted in severely truncated embryos (Yamaguchi et al., 2011). Similar truncated embryos were observed in the RNAi experiment in the present study (n=26/26). However, the phenotypes appeared to be more severe than those previously reported. Bm-wnt/RNAi embryos in this study comprised head with some epidermal structures attached. In these embryos, appendages other than the labrum and antenna were not clearly identifiable: thoracic legs were not observed, which contrasted with to the existence of immature three thoracic legs in the previous study (Fig. 5, Supplementary Fig. 5, Yamaguchi et al., 2011). Bm-ovo RNAi expression was then compared between wild type and RNAi-treated embryos fixed at about 26h AEL. In RNAi treated embryos, changes in Bm-ovo expression pattern were observed. While stripes that were clearly visible in the middle part of wild type embryos were absent in RNAi treated embryos, weak expression was noted from middle to the posterior. However, expression in anterior and posterior body region remained in treated embryos (n=19/19; Fig. 4E, F).

Discussion

Implications of the ovo homolog in Bombyx segmentation

This study revealed embryonic Bm-ovo expression patterns and some functions. Bm-ovo was dynamically expressed along the AP axis in a manner that resembled gap and pair-rule genes, and RNAi-treated embryos showed anomalies in anterior and posterior body parts, together with alterations in pair-rule gene expression patterns. The expression pattern and results of a molecular analysis of RNAi appeared to be roughly consistent with the morphological changes observed in Bm-ovo RNAi embryos. Assuming that, as a rough estimate, Bm-eve stripe #3 coincides with the posterior mesothoracic (T2) segment and the Bm-eve stripe # 2 posterior labium and Bm-run and Bm-odd stripe #2 exist in between, the weak expression of Bm-eve stripe #2 and #3 and Bm-run and Bm-odd stripe #2 may lead to aberrant gnathal/thoracic morphologies. For the observed posterior phenotypes, only the strongly expressed Bm-ovo domain appeared to be responsible. If this is the case, the abdominal phenotypes may be interpreted as the correct formation of the A1 to A5 segments with fused subsequent segments, and the region corresponding to the expression of *Bm-eve* stripe #7 contributes to the formation of the seventh abdominal segment because Bm-eve stripes#6 and #8, which strongly expressed the Bm-ovo domain cover, are expected to be mainly expressed in the posterior parts of A5 and A7, respectively.

Therefore, it is reasonable to assume that the changes observed in the expression of pair-rule genes are responsible for the observed morphological consequences. However, the mechanisms that cause these phenotypes appear to differ between the anterior and posterior regions because while the posterior phenotype was caused by deficiencies in pair-rule patterning, the pair-rule gene expression pattern in the anterior suggested that the pair-rule patterning itself is normal. The weak expression of the specific stripes of pair-rule genes is likely to be a secondary effect, i.e., caused by some mechanisms after normal patterning is achieved, such as an effect on mRNA stability.

On the other hand, given the broad expression along the AP axis of Bm-ovo, the observed phenotypes appeared to be modest; anomalies in the head or anterior abdominal segments, corresponding to strong head and weak expression around Bm-eve stripe #4 or #5, may be expected. The failure to detect corresponding phenotypes may reflect inefficiency in RNAi or the presence of redundant systems masking their manifestations.

Regulation of Bm-ovo expression

Although some irregularities were observed in segmental Bm-wntl stripes in Bm-ovo RNAi embryos, no difference was noted from wild type embryos in the posterior expression domain. Since Bm-ovo influences pair-rule gene expression as described above, the irregular stripe phenotype may be mediated through its effects on pair-rule gene functions. However, Bm-ovo does not appear to influence posterior organizing function of Bm-wnt1. On the other hand, Bm-wnt1 RNAi resulted in a change in Bm-ovo patterning. These results suggest that the regulation of Bm-ovo is controlled by the posterior organizing function of *Bm-wnt1*, appears to be complex, and may also be under the control of signaling pathways other than that of *Bm-wnt1* because *Bm-wnt1* RNAi did not result in a simple reduction in the expression of *Bm-ovo*. In *Drosophila* epidermal patterning system, ovo/svb receive inputs from Egfr pathway as well as Wingless signaling, albeit indirectly. Moreover, with its more immediate regulators, SoxN and Dichaete, ovo/svb has feedback relationships. It currently remains unclear whether similar regulatory events are operating in *Bombyx* segmentation process. However, changes in the patterning observed in Bm-ovo expression suggests a similarity in the system operating in *Bombyx* with those of *Drosophila* epidermal patterning, in which Wg represses ovo/svb expression and contributes to spatially restricting the ovo/svb expression domain. Drosophila wg is a segment polarity gene that is not involved in early segmentation events observed in short germ insects and in Bombyx. The involvement of Wg pathway in early segmentation appears to represent an ancestral condition. On the other hand, Dichaete is involved in Drosophila segmentation, with a dynamic expression pattern reminiscent of gap and pair-rule genes as in the case of Bm-ovo (Nambu and Nambu, 1996; Russell et al., 1996). The role of Dichaete in Drosophila segmentation may be an evolutionary remnant of the ancestral Wg pathway, possibly involving an *ovo* homolog. In this perspective, it is interesting that preliminary results suggested that Bombyx Dichaete (Bm-Dct) is also involved in segmentation (Supplementary Fig. 6).

Information obtained in *Drosophila* or other systems suggests the involvement of *ovo* homologs in cell differentiation and maintenance. More recently, a study of a human homolog of *ovo* indicated that it may act as a switch to enable the initiation of differentiation from the progenitor cell state by repressing stem-like properties (Renaud

et al., 2015). The involvement of *Bm-ovo* in *Bombyx* segmentation suggests the existence of a critical step common to transition from an undifferentiated to differentiated state or a cell fate decision process in ancient insect segmentation mechanisms, which is under control of *wnt* homologs.

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Figure Legends

Figure 1. *Bm-ovo* expression at early embryonic stages. Embryos were fixed at the indicated AEL times and stained with the *Bm-ovo* probe (A, C, D, F, H and I; blue) or double stained together with the *Bm-eve* probe (E, G; *Bm-ovo*: red, *Bm-eve*: blue). (B) Propidium Iodide (PI) image of A. Nuclei were not observed at this stage as shown. The anterior is to the left. (A-C) Ventral view. (D-I) Lateral view. (D-G) The ventral side is up. (H-I) The dorsal side is up. Digits indicate *Bm-eve* stripe numbers. Two weakly expressed bands are in brackets. The *Bm-ovoCp* probe was used in this experiment.

Figure 2. Morphologies of *Bm-ovo* RNAi embryos at 72 hours AEL. Wild-type (A) or *Bm-ovo* RNAi-treated embryos (B, C) were fixed at 72 hours AEL and stained with PI. Lateral view. The anterior is to the left. The dorsal side is up. Asterisks indicate prolegs. Wild-type embryos possess four proleg-bearing segments (A), whereas *Bm-ovo* RNAi-treated embryos have three (B). Note that in RNAi-treated embryos, the posterior abdominal segments and dorsal side of the meso- (T2) and meta-thoracic (T3) segments were fused (B). The dorsally fused T2 and T3 segments are in brackets. In C, irregularly shaped segments were observed (arrows). at: antenna. md: mandible. mx: maxilla. li:

labium. T1: prothorax. T2: mesothorax. T3: metathorax. A6~A9: sixth, seventh, eighth, and ninth abdominal segments, respectively.

Figure 3. Pair-rule gene expression was altered in *Bm-ovo* RNAi-treated embryos. Wild-type (A, C, E) or *Bm-ovo* RNA-treated embryos (B, D, F) were fixed at approximately 25-27 hours AEL and subjected to *in situ* hybridization in order to examine pair-rule gene expression patterns: (A, B) *Bm-eve*, (C, D) *Bm-run* and (E, F) *Bm-odd*. Lateral view. The anterior is to the left. The ventral side is up. Digits indicate the stripe number of the indicated pair-rule genes. See the text for details. The *Bm-ovoCp* probe was used in this experiment.

Figure 4. Interactions between Bm-ovo and Bm-wnt1. (A-D) Wild type and Bm-ovo RNAi treated embryos were fixed (A, B: at 23h AEL, C, D: at 26h AEL) and stained for Bm-wnt1. (E, F) Wild type and Bm-wnt1 treated embryos were fixed at 26h AEL and stained for Bm-ovo. Lateral view. The anterior is to the left. The ventral side is up. The arrow in (F) indicates the anterior boundary of the middle expression domain. See the text for details. The Bm-ovoCp probe was used in this experiment.

Figure 5. Morphologies of wild type (A) and *Bm-wnt1*RNAi treated (B) embryos. Embryos were fixed after blastokinesis (~5d AEL) and stained with PI. Lateral view. at: antenna. lr: labrum. md: mandible

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Table 1. Summary of *Bmovo* RNAi embryo phenotypes using different dsRNA.

		Observed phenotype frequency (number of observed/total					
		embryos)					
		Defects in gnathal/thoraci c segments	number of	nts			
			6	7	8		
dsRN	Bm-ov	5/14	1/14	10/14	3/14		
А	oC						
used	RNAi						
	Bm-ov	18/19	17/19	2/19	0/19		
	oE1RN						
	Ai						
	Bm-ov	11/22	16/22	6/22	0/22		
	oE3-4R						
	NAi						



Figure 1





Figure 3



Figure 4





Supplementary Figure 1. An analysis of embryonically expressed *Bm-ovo* open reading frame (ORF) species. The PCR primer pair was prepared around DNA regions corresponding to the start and stop codons in order to amplify the full ORF, and amplification was performed using cDNAs prepared at the indicated AEL times. In both cases, a strong band of approximately 2.6 kbp was detected (arrow). Other weak bands were also faintly observed; however, their identities currently remain unknown.



Supplementary Figure 2. *Bm-ovo* expression at 26h AEL stained with *Bm-ovo* probes corresponding to different regions of the *Bm-ovo* gene. Embryos were fixed and stained using the probes indicated. Lateral view. The anterior is to the left. The ventral side is up. Essentially the same staining pattern was observed for all the probes used. Scale bar: 0.2mm.



Supplementary Figure 3. *Bm-wnt1* RNAi and *Bm-ovo* RNAi were effective. Wild type, *Bm-wnt1*RNAi-, and *Bm-ovo*RNAi-treated embryos were fixed and stained with probes targeted by RNAi. (A-C) Wild type embryos (A) and embryos subjected to *Bmwnt1RNAi1*(B) and *Bmwnt1RNAi2* (C) were fixed at 26h AEL and stained with the *Bm-wnt1* probe. Wild type embryos (D) and embryos subjected to *BmovoCRNAi1*(E) were fixed at 26h AEL and stained with *Bm-ovoCp*. Wild type embryos (F) and embryos treated with *BmovoE1RNAi1*(G) and *BmovoE3-4RNAi2* (H) were fixed at 23h AEL and stained with *Bm-ovoE2p*. The anterior is to the left. (A-E) Lateral view. The ventral side is up (F-H) Ventral view. Scale bar: 0.2mm. Dashed lines approximate the boundaries of germ bands.



Supplementary Figure 4. An example of *Bm-ovo* RNAi treated embryos fixed at 23h AEL and stained for *Bm-run*. The anterior is to the left. Lateral view. The ventral side is up. Note that the intensity of #2 stripe is weaker than those of the other stripes (stripes #1, 3 and 4).



Supplementary Figure 5. Examples of mature *Bm-wnt1* RNAi treated embryos. (A) A comparison of the morphologies of a wild type (left) and *Bm-wnt1* RNAi embryo (right). In *Bm-wnt1* RNAi-treated embryos, incomplete head capsules with some epidermal structure attached posteriorly was observed. (B, B') An example of an RNAi treated embryo. Dorsal or ventral views are shown, respectively. (C, C') show another example. The anterior is to the left. As shown, gnathal appendage structures were not observed in *Bm-wnt1* treated embryos and thoracic legs were absent. Arrows indicate head capsules. Scale bar in A indicates 0.5mm.



Information on *Bombyx Dichaete (Bm-Dct)* has been deposited in KAIKObase (gene ID: BMgn003079). Based on sequence information, dsRNA targeting at *Bm-Dct* was prepared as described in the **Materials and Methods** section and subjected to an RNAi analysis. The sequences of the primers used for the amplification of the dsRNA template were: 5'-TAATACGACTCACTATAGGGAGAGATGTCGACTTTGAGCCATCAC-3', 5'-TAATACGACTCACTATAGGGAGAGGTGCCATTAGCTTGGAGAA-3'.

Supplementary Figure 6. Morphology of a *Bm-Dct* RNAi-treated embryo. Wild-type (A) or *Bm-Dct* RNAi-treated (B) embryos were fixed at 72 hours AEL and stained with PI. Although some normal morphogenesis appears to occur in the head region in the *Bm-Dct* RNAi-treated embryo, the trunk region is aberrant (n=20/20). Ir: labrum. at: antenna. md: mandible. mx: maxilla. li: labium. T1: prothorax. T2: mesothorax. T3: metathorax. Arrows indicate appendages of unknown identities.

