Recent topics on the regulatory mechanism of ecdysteroidogenesis by the prothoracic glands in insects

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INTRODUCTION

Insects are an ideal model system for studying how the regulatory mechanisms of developmental timing are coordinated with growth. The developmental transitions of insects (i.e., molting and metamorphosis) are strictly regulated by steroid hormones known as ecdysteroids. The biosynthesis of ecdysteroids, referred to as ecdysteroidogenesis, occurs predominantly in the prothoracic glands (PGs) during larval development (Gilbert et al., 2002). Since Kopeč (1922) suggested that the larval brain of the gypsy moth Lymantria dispar secretes a factor necessary for molting, attempts have been made to purify and clone this insect “brain hormone” (subsequently called prothoracicotropic hormone, PTTH). It is now widely recognized that ecdysteroid biosynthesis in the PGs is stimulated by PTTH (Kawakami et al., 1990; Mizoguchi et al., 1990); however, the importance of PTTH in the induction of molting and metamorphosis remains unclear. Removal of the brain in larvae does not result in developmental arrest, and such animals can often initiate metamorphic development, although their developments considerably delay both in the hawkmoth Manduca sexta (Truman, 1972) and in the silkworm Bombyx mori except for one hybrid race, J122 × C115 (Kobayashi, 1957). The loss of PTTH signal in the fruit fly, Drosophila melanogaster, does not cause mortality but leads to a delay in metamorphosis due to a low ecdysteroid titer, and the extended feeding period gives rise to bigger pupae and adults (McBrayer et al., 2007; Rewitz et al., 2009b). These data suggest that PTTH is required for the precise regulation of ecdysteroidogenesis but is not a crucial gatekeeper to initiate and maintain steroid biosynthesis in the PGs. In other words, other regulatory mechanisms may be involved in the regulation of ecdysteroidogenesis by the PGs.

In this review, recent studies on the regulatory mechanism of ecdysteroidogenesis by the PGs are described. In particular, I focus on humoral factors and on the neural control of ecdysteroidogenesis that have been mostly explored in B. mori.

HUMORAL FACTORS

The importance of the negative regulation of ecdysteroidogenesis has been recognized, and factors that inhibit ecdysteroidogenesis in several insect species have been described and characterized (Table 1). For example, hormonal inhibition of the PGs by the brain has been reported in locusts (Carlisle and Ellis, 1968). Crude extracts from the housefly, Musca domestica, reduce ecdysteroid synthesis and inhibit egg development when injected into the mosquito, Aedes atropalpus (Kelley et al., 1984). In the blow fly, Calliphora vicina, ecdysteroid synthesis in the larval ring gland appears to be regulated by a very potent inhibitory factor derived from the abdominal ganglion and subesophageal ganglion (Budd et al., 1993). A trypsin modulating oostatic factor (TMOF) isolated from the ovaries of adult females of flesh fly, Neobellieria bullata, is the first identified factor with prothoracicostatic activity (Hua et al., 1994). N. bullata TMOF inhibits ecdysteroid biosynthesis in the larval ring gland of flies (Hua and Koolman, 1995),

Molting and metamorphosis are strictly regulated by steroid hormones known as ecdysteroids. It is now widely recognized that ecdysteroid biosynthesis (ecdysteroidogenesis) in the prothoracic gland (PG) is regulated by the trophic factor prothoracicotropic hormone (PTTH). However, the importance of PTTH in the induction of molting and metamorphosis remains unclear, and other mechanisms are thought to be involved in the regulation of ecdysteroidogenesis by the PG. Recently, new regulatory mechanisms, prothoracicostatic factors, and neural regulation have been explored using the silkworm, Bombyx mori, and two circulating prothoracicostatic factors, prothoracicostatic peptide (PTSP) and Bommo-myosuppressin (BMS), have been identified. Whereas PTTH and BMS are secreted from the brain, PTSP is secreted from the peripheral neurosecretory system – the epiproctodeal gland – during the molting stage. The molecular basis of neural regulation of ecdysteroidogenesis has been revealed for the first time in B. mori. The innervating neurons supply both Bommo-FMRF related peptide (BRFa) and orcokinin to maintain low levels of ecdysteroids during the feeding stage. These complex regulatory mechanisms – involving tropic and static factors, peripheral neurosecretory cells as well as the central neuroendocrine system, and neural regulation in addition to circulating factors collaborate to regulate ecdysteroidogenesis. Thus, together they create the finely tuned fluctuations in ecdysteroid titers needed in the hemolymph during insect development.

Keywords: neuropeptide, prothoracicostatic factor, innervating neuron, prothoracic gland, Bombyx mori, ecdysteroidogenesis, peripheral neurosecretory cell

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but *N. bullata* TMOF is not present in the central nervous system of the larvae (Bylemans et al., 1996). Allatostatin-B (AST-B) isolated from adult brains of the cricket, *Gryllus bimaculatus*, has been observed to decrease the hemolymph ecdysteroid titer in the adult of *G. bimaculatus* (Lorenz et al., 1998) but the PGs do not produce physiologically significant amounts of ecdysteroids at this stage (Hoffmann et al., 1998). Thus, no endogenous prothoracicostatic factor had been identified until the isolation of prothoracicostatic peptide (PTSP) from larval brains of *B. mori* (Hua et al., 1999).

Prothoracicostatic peptide has the same sequence as Manse-myoinhibitory peptide-I (MIP-I), which was previously isolated from the nerve cord of the adult *M. sexta* (Blackburn et al., 1995). PTSP also shows high sequence homology to *G. bimaculatus* AST-B. This peptide inhibits both basal and PTTH-stimulated ecdysteroidogenesis in *vitro* in a dose-dependent manner, and injection of synthetic PTSP has been shown to temporally decrease the hemolymph ecdysteroid titer in fifth instar larvae of *B. mori* (Liu et al., 2004). Interestingly, the peripheral neuroendocrine system – the epiproctodeal gland – is the principal site of neurohemal release of MIP in *M. sexta* (Davis et al., 2003): MIP-immunostaining in the epiproctodeal gland begins to increase during the first half of the molting period, and then most of the immunostaining is rapidly lost within several hours. This rapid loss of immunostaining suggests that a massive release of MIP from the epiproctodeal gland into hemolymph occurs at this stage, when there is a rapid decline in the ecdysteroid titer. The release of PTSP from the epiproctodeal gland also appears to occur at the same stage in *B. mori*, while the expression of *ptsp* in the brain is high during the feeding stage and low during the molting stage (Yamanaka et al., 2010). Recently, a *B. mori* G protein-coupled receptor, which had previously been identified as an ortholog of the *Drosophila* sex peptide receptor (Yapici et al., 2008), is characterized as a functional PTSP receptor (Yamanaka et al., 2010). This receptor responds specifically to PTSP when examined using a heterologous expression

### Table 1 | List of peptides and biogenic amine taken up in this review.

| Name                                      | Structure                              | Tropic/static | References             |
|-------------------------------------------|----------------------------------------|---------------|------------------------|
| **B. MORI**                               |                                        |               |                        |
| Prothoracicotropic hormone (PTTH)         | GNIQVENAIPOPPCTKYKKEIEDLGENSVPRFIET   | Tropic        | Kavakami et al. (1990) |
|                                           | RNCNKTQOPTCRPFCYKCESLYSITILKRETQGUESL |               |                        |
|                                           | EIPELKYRWVXASHPVSAACCLCRDYLQRNYNN     |               |                        |
| **PROTHORACICOSTATIC PEPTIDES (PTSPs)**   |                                        |               |                        |
| PTSP-I                                    | AWQDLNSAWa                             | Static        | Hua et al. (1999), Yamanaka et al. (2010) |
| PTSP-II                                   | GWQDLNSAWa                             |               |                        |
| PTSP-III                                  | APEKWAFHGSWa                           |               |                        |
| PTSP-IV                                   | GWNDISSWVa                             |               |                        |
| PTSP-V                                    | AWQDMSAWa                              |               |                        |
| PTSP-VI                                   | AWSALHTWa                              |               |                        |
| PTSP-VII                                  | AWQDLNSWA                               |               |                        |
| PTSP-VIII                                 | AWSLHSGWa                              |               |                        |
| **BOMMO-MYOSUPPRESSIN (BMS)**             |                                        |               |                        |
| **BOMMO-FMRF-AMIDE-RELATED PEPTIDE (BRFa)** |                                      |               |                        |
| BRFa-I                                    | SAIDRSMIRFa                            | Static        | Yamamoto et al. (2006) |
| BRFa-II                                   | SASFVRFa                               |               |                        |
| BRFa-III                                  | DPSFIFRa                               |               |                        |
| BRFa-IV                                   | ARNHFRLa                               |               |                        |
| **ORCOKININ**                             |                                        |               |                        |
| Bommo-orc-I                               | NFDEIDESSLNTFV                         | Tropic        | Roller et al. (2008), Yamanaka et al. (2011) |
| Bommo-orc-II                              | NFDEIDRSMFFPYAI                        |               |                        |
| **FXPRL-AMIDE**                           |                                        |               |                        |
| Diapause hormone (DH)                     | TDMKDESDRGASGERALWFGPRLa               | Tropic        | Watanabe et al. (2007) |
| **BIOGENIC AMINE**                        |                                        |               |                        |
| Octopamine                                 |                                        |               |                        |
| **OTHER INSECTS**                         |                                        |               |                        |
| *N. bullata* TMOF                         | NPTNLH                                 | Static        | Hua et al. (1994), Hua and Koolman (1995) |
| *G. bimaculatus* AST-B                    | GWQDLNGGWa                             | Static        | Lorenz et al. (1998)   |
| *H. armigera* DH-like peptide             | NDVKDGAASGAHSDLGLWFGPRLa               | Tropic        | Zhang et al. (2004)    |
| *N. bullata* pyrokinin-II                 | SVQFKPRLa                              | Tropic        | Verleyen et al. (2004) |

## Notes

- Allatostatin-B (AST-B) was isolated from adult brains of the cricket, *Gryllus bimaculatus*.
- PTSP inhibits basal and PTTH-stimulated ecdysteroidogenesis in *vitro*.
- PTSP has been shown to decrease the hemolymph ecdysteroid titer in fifth instar larvae of *B. mori*.
- MIP-immunostaining in the epiproctodeal gland increases during the first half of the molting period and rapidly declines.
- A G protein-coupled receptor for PTSP has been identified in *B. mori*.
system. The receptor is expressed at low levels during the feeding stage, but is highly expressed in the PGs on the day before each larval and pupal ecdysis, when massive release of PTSP from the epiproctodeal gland occurs. These results suggest that PTSP secreted from the epiproctodeal gland acts on the PGs through the PTST receptor during the molting cycle in B. mori and that the peripheral neurosecretory cells, as well as the neurosecretory system in the central nervous system, play a stage-specific role in regulating ecdysteroidogenesis. PTSP is unlikely to be involved in initiating the decrease in the ecdysteroid titer, but is probably involved in either full inactivation or suppression of re-activation of the PGs to finely regulate the ecdysteroid titer in the hemolymph (Yamanaka et al., 2010).

Another prothoracicostatic factor purified from pupal brains of B. mori is a decapeptide that shows the conserved structure of an insect myosuppressin, and has thus been named Bommo-myosuppressin (BMS; Yamanaka et al., 2005). Myosuppressin is a member of a larger peptide family of FMRF-amide-related peptides (FaRPs), which are known to regulate a wide range of processes from behavior to physiology in invertebrates (Mackey et al., 1987; Nelson et al., 1998; Nicols, 2003). BMS dose-dependently suppresses the cAMP level and inhibits ecdysteroidogenesis in the larval PGs at much lower concentrations than PTSP. The specific receptor for BMS is highly expressed in the PGs at the feeding stage. Together, these results suggest that BMS is secreted from the brain and plays an important role in maintaining low ecdysteroid levels during the feeding stage in the hemolymph. Vertebreates including humans have also been shown to possess peptides with RF-amide at the C-terminus (Yang et al., 1985), and at least one RF-amide peptide has been shown to inhibit steroid hormone release in experiments using an in vitro adrenal slice culture system (Labrouche et al., 1998). RF-amide peptides may modulate steroid hormone secretion throughout metazoans, thereby controlling post-embryonic development.

FXPRL-amide peptides, which are not homologous with PTTH, can stimulate ecdysteroidogenesis by the PGs. Diapause hormone (DH) which was identified as an inducer of embryonic diapause in B. mori (Imai et al., 1991), stimulates ecdysteroidogenesis by the PGs of late fifth instar larvae of B. mori (Watanabe et al., 2007). The receptor for DH is expressed in the PGs, as well as in pupal pheromone glands and ovaries prior to eclosion, and the expression of this receptor in the PGs increases from the last half of the fifth instar stage to the first day of the pupal stage. These results suggest that DH regulates ecdysteroidogenesis during the late stages of fifth instar larvae in B. mori. Possible prothoracicotropic effects of FXPRL-amide peptides have been reported in other insects. In the cotton bollworm Helicoverpa armigera, DH-like peptide can break the pupal diapause by activating the ecdysteroidogenesis by PGs in a temperature-dependent manner (Zhang et al., 2004). The pupariation of N. bullata, is accelerated by a FXPRL-amide peptide, N. bullata pyrokinin-II (Verleyen et al., 2004). Taken together, these results suggest that FXPRL-amide peptides can function as prothoracicotropic factors across insect species and may play important roles in post-embryonic insect development.

NEURAL REGULATION

Since the first report of neural innervation in the PGs by Lyonet (1762; Figure 1A), the importance of PG-innervating neurons in the control of ecdysteroidogenesis has been well documented (Scharrer, 1964; Srivastava and Singh, 1968; Hintze-Podulaf, 1970; Giebultowicz and Denlinger, 1985). In B. mori, the PGs are innervated by seven nerves starting from the subesophageal ganglion, prothoracic ganglia and mesothoracic ganglia, and by unpaired nerves between the above mentioned ganglia (Figure 1B, Yokoyama, 1956). A positive correlation between the levels of electrical activity of PG-innervating neurons and the ecdysteroid titer during the course of development in the last larval instar has been reported in the American cockroach Periplaneta americana (Richter and Gersch, 1983), while the electrical activity shows an inverse relationship with ecdysteroid titer in the cabbage moth Mamestra brassicae (Okajima and Watanabe, 1989).
However, studies revealing the molecular basis of regulation by the PGs have been restricted to hormonal factors throughout the last century.

Recently, for the first time we revealed the molecular basis of neural regulation of ecdysteroidogenesis in *B. mori* (Yamanaka et al., 2006). We found that Bommo-FMRF related peptide (BRFa) regulates ecdysteroidogenesis through direct innervation of the PGs in *B. mori*. BRFa is predominantly expressed in the prothoracic ganglion, and neurons in the prothoracic ganglion innervate the PGs and supply BRFAs to the surface of PGs (Figure 1B). Electrophysiological recordings during development confirmed the increased firing activity of BRFa-expressing neurons in developmental stages of low PG activity and decreased ecdysteroid levels in the hemolymph. The prothoracotropic function of BRFa has been confirmed by the report that BRFa neurons suppress the expression of several P450 genes induced by PTTH in the PGs (Yamanaka et al., 2007).

We found that other neuropeptides are delivered to the surface of the PGs through PG-innervating neurons (Yamanaka et al., 2011). Using direct mass spectrometric profiling of axons associated with the PGs, we detected several peptide peaks corresponding to orcokinin gene products in addition to BRFAs. One of the two pairs of BRFa-expressing neurosecretory cells in the prothoracic ganglion co-expresses orcokinin, and these neurons project axons through the transverse nerve and terminate on the surface of the PGs (Figure 1B). Orcokinin was originally isolated from the nervous system of the spinycheek crayfish *Orconectes limosus* as a myostimulatory factor (Stangier et al., 1992). Orcokinin-like peptides have subsequently been identified in several insects (Pascual et al., 2004; Hofer et al., 2005; Liu et al., 2006; Christie, 2008; Roller et al., 2008; Clynen and Schoofs, 2009), although their physiological functions remain largely unknown except for a few cases (Hofer and Homberg, 2006). In *B. mori*, orcokinins have clear prothoracotropic activity and are able to cancel the static effect of BRFa on ecdysteroid biosynthesis, whereas the suppressive effect of BRFa on cAMP production remains unchanged in the presence of orcokinins (Yamanaka et al., 2011). The acute effects of prothoracicotropic or prothoracostatic factors which becomes manifest temporally during short-term *in vitro* incubation of the PG are mediated by translation and phosphorylation of intercellular molecules involved in the signal transduction of ecdysteroidogenesis (Gilbert et al., 2002; Huang et al., 2008; Rewitz et al., 2009a). On the other hand, the chronic effects of these factors which become obvious on a longer time scale during development are mediated by the transcriptional regulation of some genes encoding ecdysteroidogenenic enzymes probably via cAMP-mediated pathway (Gilbert et al., 2002; Huang et al., 2008). The chronic effects of these factors are pharmacologically separable from the acute effects (Yamanaka et al., 2007). Thus, orcokinins might regulate ecdysteroidogenesis by the PGs via cAMP-independent way; that is, they specifically eliminate the acute prothoracostatic effect of BRFAs while maintaining the chronic inhibitory effect of BRFAs. The signaling pathway of orcokinin at the molecular level needs to be elucidated to prove this hypothesis.

What is the functional significance of such a complex feature of the PG-innervating neurons in *B. mori*? It is known that a low level of ecdysteroid titer during the intermolt period has a complex effect on the regulation of tissue growth and proliferation (Champlin and Truman, 1998; Colombani et al., 2005; Nijhout et al., 2007). Therefore, orcokinins might maintain a low ecdysteroid titer while BRFa suppresses full potentiation of the PGs prior to the ecdysteroid pulse generated by PTTH (Mizoguchi et al., 2001).

Prothoracic gland-innervating neurons of other lepidopteran insects are also stained by both FMRFa- and orcokinin-specific antibodies (Tanaka, unpublished data).
of bilaterally symmetric neurons located in the cerebral labrum portion of the brain directly innervate the PGs (Sigmund and Korge, 2001) and induce the production and secretion of ecdysone by delivering PTTH directly (McBrayer et al., 2007); however, it is not certain whether other peptides are delivered to the PGs through these innervating neurons. By contrast, PG-innervating neurons are not stained by PTTH-specific antibody in B. mori (Tanaka, unpublished data). Different peptidergic systems may regulate ecdysteroidogenesis among the various insect species.

FUTURE PROSPECTS

Bombyx mori has been a useful tool for insect endocrinology since the discovery of PG function by Fukuda (1944), and studies have shown that ecdysteroidogenesis by the PGs is under the control of a complex mechanism (Figure 2). Tropic and static factors, the peripheral neurosecretory system as well as the central neuroendocrine system, and neural regulation in addition to circulating factors all collaborate to regulate ecdysteroidogenesis by the PGs. Thus, together they create the finely tuned fluctuations in ecdysteroid titer needed in the hemolymph during the development of B. mori. Moreover, we should carefully re-investigate factors and mechanisms that have been ignored for a long time. For example, the roles of biogenic amines in ecdysteroidogenesis remain unclear, although it has been reported that octopamine stimulates the production of ecdysteroids by larval PGs of B. mori in vitro (Hirashima et al., 1999). How the PGs relay their condition to the neural system is not yet fully understood, but the PG-innervating neurons may be involved in this communication. Three PG-innervating neurons contain afferent neurons as well as efferent ones in M. brassicae (Okajima and Watanabe, 1989). Although the function of these afferent neurons is unknown, the CNS may be monitoring the condition of the PGs via these neurons.

The neuropeptides responsible for the ecdysteroidogenesis in B. mori are commonly present among insect species, but their function in ecdysteroidogenesis remains to be elucidated in other insects. No prothoracicostatic factor has been identified as yet in D. melanogaster, and the importance of insulin signaling in regulating ecdysteroid production by the PGs has been recently highlighted (see the review by Marchal et al., 2010 and the recent topics by Gibbens et al., 2011). The regulatory mechanisms of ecdysteroidogenesis by the PGs may be diverse among insect species and need to be elucidated in different insect models in order to understand how the insects adapted to diverse environment.

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