The EGPs: The Eclosion Hormone and Cyclic GMP-Regulated Phosphoproteins. I. Appearance and Partial Characterization in the CNS of Manduca sexta

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The present study describes 2 phosphoproteins, both with an apparent molecular weight of 54 kDa, in the CNS of the tobacco hornworm, Manduca sexta. Their phosphorylation is regulated by a neuropeptide, eclosion hormone (EH), and the second messenger cGMP, which thus have been named the EGPs (eclosion hormone- and cGMP-regulated phosphoproteins). Although cAMP was more effective than cGMP at stimulating the phosphorylation of the EGPs in CNS homogenates, in the intact CNS cGMP was more effective. Since cGMP mediates the action of EH, this strongly suggests that cGMP is the second messenger that stimulates the phosphorylation of the EGPs in vivo.

The EGPs can only be phosphorylated in vitro during discrete time periods during the life of Manduca. During the larval and pupal molts, the EGPs can first be phosphorylated just prior to ecdysis. Their ability to be phosphorylated is correlated with the time when the insect is sensitive to EH. This close temporal correlation suggests that the ability to phosphorylate the EGPs determines when the insect can first respond to EH. During adult development, the EGPs first appeared on fluorograms 6 d before sensitivity to EH, suggesting that at this stage other factors may also be involved in the regulation of sensitivity. For the ecystes of all 3 stages, EH appeared to stimulate the phosphorylation of the EGPs at ecdysis.

The EGPs were found in all regions of the prepupal nervous system that were investigated, but only in the abdominal and pterothoracic ganglia of the developing adult. Fractionation of nervous system homogenates by ultracentrifugation revealed that one of the EGPs was present only in the pellet fraction, whereas the other was approximately equally distributed between pellet and supernatant. Furthermore, the EGPs in the pellet fraction could be partially solubilized with detergents and high salt concentrations.

In eukaryotic cells, the second messengers cAMP and cGMP exert most of their effects through the phosphorylation of specific proteins (see Nestler and Greengard, 1984). In the nervous systems of both vertebrates and invertebrates, many neurotransmitters, neuromodulators, and hormones exert their effects through the elevation of cyclic nucleotides (Lingle et al., 1982; Drummond, 1984) and, hence, through the phosphorylation of particular proteins (Nestler and Greengard, 1984).

In insects, the neuropeptide, eclosion hormone (EH), acts directly on the CNS to release the motor programs for the stereotyped ecdysis behavior (Truman, 1978a) and to activate stage-specific reflexes (Levine and Truman, 1983). These actions of EH are mediated through an increase in the intracellular levels of cGMP (Morton and Truman, 1985). Studies of EH action during pupal ecdysis of Manduca subsequently identified 2 proteins, both with an apparent molecular weight of 54,000 Da, which are phosphorylated in response to EH (Morton and Truman, 1986). These proteins were initially called "the 54 kDa proteins," but are now referred to as the EGPs—the eclosion hormone- and cGMP-regulated phosphoproteins.

EH is released periodically during the life of Manduca to trigger the ecysts of larval, pupal, and adult stages. In each case the nervous system is sensitive to the peptide only during the final few hours preceding ecdysis. Responsiveness is then lost at ecdysis and is only regained as the insect is undergoing the next molt. The study of pupal ecdysis showed a close correlation between the ability to phosphorylate the EGPs in vitro and the development of sensitivity to EH (Morton and Truman, 1986). The present paper further explores the relationship of the EGPs to EH action during pupal ecdysis and also at the larval and adult stages of the insect's life history.

Materials and Methods

Rearing and staging of animals. Larvae of the tobacco hornworm Manduca sexta were reared individually on an artificial diet (Bell and Joachim, 1978) at 26°C under a 17L:7D photoperiod. At the end of the terminal (fifth) larval stage the insect ceased to feed, emptied its gut, and entered the "wandering" stage. Pupal ecdysis occurred 4 d later. Once adult development was initiated, the animals were kept at 26°C with a 12L:12D photoperiod, with lights-off arbitrarily designated as 24:00. Adult ecysts occurred 18-20 d later.

The timing of ecdysis of the fifth larval stage was predicted using external morphological markers, as described by Copenhaver and Truman (1982). These were the first appearance of pigmentation in the new mandibles (10 hr before ecdysis) and the appearance of air in the old head capsule due to reabsorption of the molting fluid (6 hr before ecdysis). Events prior to pupal ecdysis were timed with reference to the pigmentation of a pair of sclerotized bars on the dorsal surface of the metathoracic segment (24 hr before ecdysis) and to the resorption of molting fluid in the anterior segments (4 hr before ecdysis; Truman et al., 1980). Animals at these stages will subsequently be referred to as -24 hr and -4 hr animals, respectively. As the timing of adult ecdysis is controlled by a circadian clock (Truman, 1978b), animals were taken...
at various times after “lights-on” (12:00) on the last day of adult development, with ecdysis occurring between 21:00 and 00:30 (Reynolds et al., 1979). For each experiment, parallel groups of animals were allowed to ecdyse and the timing of the experimental animals was judged relative to that of these controls.

**Endogenous phosphorylation of homogenates.** Nervous tissue was removed and homogenized in 50 mM HEPES buffer, 5 mM EDTA, 1 mM dithiothreitol (DTT), pH 7.0, at a concentration of 1–1.5 mg protein/ml of buffer. The phosphorylation was carried out by incubating 100 μl of the homogenate with 40 μl 250 mM HEPES buffer, pH 7.0, 20 μl 100 mM MgCl₂, 20 μl 10⁻⁵ M ATP containing 10 μCi gamma-³²-P-ATP (New England Nuclear; approximately 3000 Ci/mmol), and 20 μl cGMP, cAMP, or water. The reaction was carried out at 30°C for 5 min and was started by the addition of the ATP and stopped with 200 μl 20% trichloroacetic acid (TCA). The reaction mixture was allowed to stand on ice for 30 min, centrifuged at 10,000 × g for 5 min at 4°C, and the supernatant separated by 2-dimensional SDS-PAGE, as described below.

**Back-phosphorylation of the isolated CNS.** Abdominal nervous systems were dissected, leaving as much of the tracheal supply intact as possible, and rinsed in Grace’s insect medium (Gibco). The nervous systems were then placed in 0.5 ml of Grace’s medium in the presence or absence of EH and incubated at 26°C in a 95% O₂-5% CO₂ atmosphere. Tissues were removed at the allotted time, homogenized and phosphorylated in vitro in the presence of 0.1 mM cGMP and ³²P-ATP, as described above. Any proteins phosphorylated as a result of incubation with EH would then be unavailable for phosphorylation by labeled phosphate and would be absent on the fluorograms.

The EH was partially purified from the corpora cardiaca–corpora allata complex of pharate adult moths by passing an acidified methanol extract through a C-18 reverse-phase column using high-performance liquid chromatography. The concentration of EH was measured by bioassay, as described by Truman et al. (1980), immediately before use.

**Separation and visualization of the proteins.** The first dimension of separation was by nonequilibrium pH gradient electrophoresis (NEPHGE, O’Farrell et al., 1977), with gels containing 9.2 mM urea, 4% acrylamide, 2% Nonidet P-40, and 2% Ampholines, pH 3.5–10 (both LKB), 5% 2-mercaptoethanol (2-ME), and 0.01% SDS. The samples were left at room temperature overnight, vortexed, and centrifuged at 10,000 × g for 5 min at 4°C and the supernatant separated by 2-dimensional SDS-PAGE, as described below.

**Photophosphorylation of the isolated CNS.** Homogenates of the CNS were separated into soluble and particulate fractions to determine the location of the EGPs. Homogenates were phosphorylated (at 3 times normal volumes), the reaction stopped by cooling on ice, and then were centrifuged at 100,000 × g for 1 hr. The protein in the supernatant was precipitated by the addition of 600 μl 20% TCA, and the pellet fraction dissolved in 40 μl sample buffer. The proteins were separated by 2-dimensional SDS PAGE, as described above.

**Photoaffinity labeling of cAMP-binding proteins.** cAMP-binding proteins present in homogenates were identified by photoaffinity labeling with 8-azido cAMP according to the methods of Walter and Greengard (1983). Nervous tissue was homogenized in 50 mM 2-(N-morpholino)ethanesulfonic acid (MES) buffer, pH 6.2, containing 1 mM IBMX, 10 mM MgCl₂, and 0.5 mM 2-ME, at a concentration of about 1 mg/ml and 80 μl, incubated with 10 μl of 10⁻⁷ M 8-azido cAMP (ICN, 71 Ci/mmol) in the presence or absence of 10 μl of 5 × 10⁻⁵ M cAMP, both dissolved in the same buffer for 1 hr in the dark. The samples were then exposed to UV light at 254 nm using a Minirell UVG-11 lamp (Ultraviolet Products) placed 10 cm above the samples. The reaction was stopped and the proteins separated by 2-dimensional SDS-PAGE.

**Results**

**Phosphorylation of the EGPs in CNS homogenates in the presence of cGMP and cAMP**

Figure 1.4 gives an example of a portion of a fluorogram that shows the phosphorylation of the 2 EGPs in response to the addition of 0.1 mM cGMP to a cell-free CNS homogenate. The incorporation of phosphate into the EGPs was estimated by measuring the density of the EGP-A. The path used in the scan also measured the density of a 40 kDa protein whose phosphorylation was cyclic nucleotide-independent, and of a 36 kDa protein whose phosphorylation was dependent on cyclic nucleotide levels (Fig. 1D). As the incorporation of label into the 40 kDa protein was not cyclic nucleotide-dependent, it was used as a standard for comparing the incorporation of label into the EGPs.

The effect of various concentrations of cAMP and cGMP on the incorporation of labeled phosphate into EGP-A is shown in Figure 2. cAMP was more effective than cGMP at stimulating the phosphorylation of the EGPs in CNS homogenates. The concentration of cAMP needed to stimulate half-maximal phosphorylation of the EGPs was 0.2 μM, compared to 36 μM for cGMP. Both cyclic nucleotides stimulated the same maximal level of phosphorylation. Other cyclic nucleotide-stimulated phosphoproteins, such as the 36 kDa protein, showed a similar dose–response relationship, with cAMP being more effective than cGMP.

**Time course of EGP phosphorylation in the isolated CNS**

In a previous report (Morton and Truman, 1986) we showed, by using the back-phosphorylation technique of Forn and Greengard (1978), that the injection of EH into prepupa stimulated the phosphorylation of the EGPs. To show that this phos-
Figure 1. Estimation of the incorporation of label into the EGPs. A, Portion of a fluorogram showing the region scanned with the densitometer. In this and subsequent figures, the portion of the fluorogram shown covers the molecular-weight range of about 70–30 kDa and the isoelectric point range of 5.0 (left) and 7.0 (right). The 2 EGPs are marked with arrows and the density of the more acidic EGP (EGP-A) measured relative to the standard phosphoprotein at 40 kDa, which is marked with an asterisk. B, Examples of densitometer scans from fluorograms made from abdominal nervous system homogenates incubated in the presence or absence of 100 μM cGMP. The peaks corresponding to EGP-A, the 40 kDa standard phosphoprotein and the 36 kDa phosphoprotein are marked with arrows. The area of the EGP peak was then measured relative to the area of the 40 kDa peak.

Phosphorylation was due to the direct action of EH on the CNS, we removed the abdominal nervous systems from -4 hr animals and incubated them for 1 hr in the presence or absence of 1 U/ml of EH. After incubation, the nervous tissue from the 2 groups was homogenized and incubated with 32P-ATP in the presence of 0.1 mM cGMP. Nervous tissue incubated in the presence of EH showed no incorporation of label into the EGPs (Fig. 3). Controls incubated without hormone showed the normal phosphorylation patterns. The lack of label in the former was presumed to be the result of the EGPs' being phosphorylated
with unlabeled phosphate in response to EH exposure. The EGPs were the only proteins that consistently showed a lack of labeling after intact nervous tissue was incubated in the presence of EH. For example, in Figure 3.4 the 36 kDa protein (asterisk) showed the same level of incorporation whether or not the CNS was exposed to EH.

The specificity of the action of EH in phosphorylating only the EGPs was also seen when nervous tissue from 24 hr animals was incubated in the presence or absence of EH. At this stage the EGPs are presumed absent from the CNS (Morton and Truman, 1986), but EH will elevate cGMP levels (Morton and Truman, 1985). Inspection of fluorograms showed no detectable changes in the incorporation of labeled phosphate into proteins from these samples that resulted from exposure to EH (results not shown).

At pupal ecdysis, when ~4 hr animals are injected with EH, the latency to ecdysis is inversely related to hormone dosage, with a minimum latency of about 50 min (Truman et al., 1980). Prepupae injected with a maximal dosage of EH, however, show an increase in cGMP within 5 min (Morton and Truman, 1985). When nervous tissue is incubated in a high concentration of EH (1 U/ml), the EGPs are completely phosphorylated by 60 min (Fig. 3.4). At lower hormone concentrations (0.25 U/ml), the time taken for phosphorylation of the EGPs is longer (as measured by the reduced incorporation of labeled phosphate into proteins; Fig. 3B). Thus, as is seen with the behavioral latency, the rate at which the EGPs are phosphorylated also appears to be a function of the amount of EH that is added.

The use of the isolated abdominal CNS also enabled us to test the relative abilities of cAMP and cGMP to stimulate the phosphorylation of the EGPs in the intact CNS. Abdominal nervous systems were incubated in various concentrations of cGMP and CAMP for 1 hr and then subjected to in vitro phosphorylation, as above. At 1 mM, both cGMP and CAMP stimulated complete phosphorylation of the EGPs, so that no labeled phosphate was incorporated into the EGPs during the subsequent in vitro phosphorylation (Fig. 4). At lower concentrations, cGMP was consistently more effective than cAMP in stimulating the endogenous phosphorylation of the EGPs, with cGMP showing an ED50 of about 0.1 µM, whereas that of cAMP was 4 µM. Thus, although cAMP was more effective than CAMP at stimulating the phosphorylation of the EGPs in cell-free homogenates, the reverse relationship was seen when the intact CNS was exposed to the cyclic nucleotides.

The EGPs were the only proteins detected whose phosphorylation was stimulated more effectively by cGMP than by CAMP in the intact CNS. The 36 kDa protein was also phosphorylated in the isolated CNS in response to cyclic nucleotides. Unlike the case with the EGPs, however, cAMP was more effective than cGMP in stimulating its phosphorylation. Incubation of nervous systems for 1 hr with 1 mM cAMP resulted in a reduction of 51 ± 17% (mean ± SEM; n = 3) in the incorporation of label into the 36 kDa protein during subsequent in vitro phosphorylation, whereas the same levels of cGMP were ineffective (reduction of 7 ± 10%).

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Temporal pattern of appearance of the EGPs during the life history of Manduca

EH triggers larval, pupal, and adult ecdyses in Manduca (Truman, 1971; Truman et al., 1980; Copenhaver and Truman, 1982). If the EGPs are involved in the action of EH in triggering ecdysis, then they should be present prior to each ecdysis, irrespective of the stage in the insect's life history.

Figure 5A shows the temporal relationship between the incorporation of labeled phosphate into the EGPs in the abdominal nervous system and the behavioral sensitivity to EH during the fifth larval and pupal molts. During the last larval molt, behavioral sensitivity to EH begins at about 6 hr prior to ecdysis (Copenhaver and Truman, 1982). When abdominal nervous systems were taken from molting larvae at this stage and subjected to in vitro phosphorylation, we found a very low level of incorporation of label into the EGPs. Two hours later, the EGPs were clearly labeled and, after another 2 hr, the density was even greater. Approximately 2 hr later, the larvae had released EH and were undergoing ecdysis; extracts of the nervous tissue taken at this time no longer showed incorporation of phosphate into the EGPs. This abrupt disappearance of the ability to label these proteins with exogenous phosphate was presumed to be due to endogenous EH’s having stimulated their phosphorylation with endogenous unlabeled phosphate.

Following ecdysis, no incorporation of labeled phosphate into the EGPs was detected during the feeding stage of the fifth instar, the wandering stage, and most of the prepupal period (a period of 7 d). The absence of labeling of the EGPs was correlated with the fact that insects are not responsive to EH during this period. Both behavioral sensitivity and the ability to label the EGPs reappeared 8 hr before pupal ecdysis (Morton and Truman, 1986). As in larval ecdysis, the onset of pupal ecdysis behavior
Figure 3. Back-phosphorylation of the EGPs in the intact, isolated CNS by EH. A, Example of fluorograms made from the abdominal nervous systems from -4 hr prepupae that were incubated in culture in the presence or absence of 1 U/ml of EH for 1 hr. After incubation, the nervous systems were homogenized (2 nervous systems per sample) and phosphorylated with 32P-ATP in the presence of 0.1 mM cGMP. The proteins were separated by 2D SDS-PAGE and fluorograms made. The 2 EGPs are marked with arrows in the control fluorogram (i; incubated in the absence of EH), but are absent after incubation in 1 U/ml of EH (ii). B, Time course of phosphorylation of the EGPs in nervous systems that were exposed to 0.25 units of EH/ml for various intervals before homogenization and in vitro phosphorylation in the presence of 0.1 mM cGMP. The incorporation of labeled phosphate into the EGPs was estimated as described in Figure 1. Controls represent nervous systems that were incubated in the absence of EH for 2 hr. Each time point represents data from one sample (2 nervous systems), except for the control and 0 time points, which represent the mean and range of 2 samples.

marked the abrupt disappearance of the ability to incorporate phosphate into the EGPs.

We encountered some difficulty in reliably detecting the EGPs in the abdominal nervous system of pharate adult Manduca. This was because this tissue contained more protein at this stage (210 μg, compared to 176 μg for prepupae) because of the presence of the “dorsal pad” of connective tissue. The presence of another phosphoprotein at about 57 kDa partially obscured the
EGPs and made it difficult to reliably detect their presence (see Fig. 6). To partially overcome this problem we phosphorylated abdominal nervous system homogenates and then centrifuged them at 100,000 × g for 1 hr. The proteins present in the supernatant were then separated by SDS-PAGE. As detailed below for the prepupal CNS, only EG-P-A was present in the supernatant fraction, and this was used as a marker for the presence of both EGPs at this stage.

The time course of the appearance of EH sensitivity and the ability to label EG-P-A in the abdominal nervous system of the developing adult are shown in Figure 5B. The presence of the 57 kDa protein made densitometric quantification of EG-P-A during adult development unreliable (see Fig. 6), so the level of incorporation of label into EG-P-A at various times was judged subjectively and plotted relative to the incorporation seen at 3 hr before ecdysis. Behavioral sensitivity to EH was first clearly seen between 6 and 8 hr before ecdysis. Abdominal nervous systems taken from animals at this time and at times closer to ecdysis showed the presence of EG-P-A. As with larval and pupal ecdysis, abdominal nervous systems removed from animals just after adult ecdysis no longer showed incorporation of label into EG-P-A. This suggests that EH also stimulated the phosphorylation of the EGPs at adult ecdysis. Unlike in larvae and pupae, however, incorporation of label into EG-P-A was clearly seen at times when developing adults were not behaviorally sensitive to EH. They could first be labeled as early as 6 d before ecdysis. Another unusual feature of adult ecdysis was that at 24 hr after ecdysis, EG-P-A was again able to incorporate label.

Figure 6 shows the results of an experiment to determine whether EH treatment of developing adults stimulates the phosphorylation of EG-P-A at times when EH will not trigger ecdysis behavior. Animals at 28 hr before ecdysis will not respond to EH, whereas 6 hr before ecdysis they will respond to EH by initiating ecdysis 3.5 hr after injection. Animals were injected at various times with EH, and the percentage of animals that showed premature ecdysis behavior noted. Data redrawn from Copenhaver and Truman (1982), Morton and Truman (1986), and Reynolds et al. (1979). Lower panels represent the presence of the EGPs during development. Abdominal nervous systems were removed and homogenized at various times during development, phosphorylated with ³²P-ATP in the presence of 0.1 mM cGMP, and the proteins separated by 2D SDS-PAGE. Fluorograms of the gels were made and the incorporation of labeled phosphate into the EGPs was estimated as described in Figure 1 for larval and pupal ecdysis. For adult ecdysis, the relative incorporation of label was judged subjectively relative to the incorporation seen 3 hr before ecdysis. A score of 0 indicates that no incorporation of label into the EGPs could be detected, while a score of 1 indicates that incorporation was approximately equal to that seen at −3 hr. A, Larval and pupal development. Each day of development is marked relative to 3 developmental markers: the ecdysis into the fifth larval instar (V), wandering (W), and pupal ecdysis (P). B, Adult development. Each time point represents either days after pupal ecdysis (P) or days before or after adult ecdysis (D).
system and also in the fused pterothoracic ganglion (total homogenates). In pterothoracic ganglia removed from moths that had just eclosed, as in the abdominal nervous system, the EGPs no longer accepted labeled phosphate. The EGPs were also seen in the pterothoracic ganglion 24 hr before ecidyse, when the animals were not behaviorally sensitive to EH. We found no definitive evidence for the presence of the EGPs in the pterothoracic ganglion or the brain. This inability was partly due to the presence of a 57 kDa protein, similar to the one found in the abdominal nervous system, which may have obscured low levels of labeling in the 54 kDa region. In some fluorograms, very faint labeling was seen in the 54 kDa region, but we could not be confident as to whether this was due to low levels of the EGPs or to smearing of the 57 kDa protein.

**Distribution of the EGPs in subcellular fractions**

Figure 7 shows the distribution of the EGPs in the supernatant and pellet fractions after centrifugation at 100,000 × g. Homogenates of abdominal nervous tissue from −4 hr prepupae, which had been phosphorylated and then centrifuged, EGP-A was found in both the soluble and particulate fractions, whereas the more basic EGP (EGP-B) was found only in the particulate fraction (Fig. 7, A, B). In one series of experiments the homogenate was centrifuged before phosphorylation, the pellet resuspended in homogenization buffer, and both fractions phosphorylated independently in the presence of 0.1 mM cGMP. No label was incorporated into the EGPs in the pellet fraction, presumably because the *Manduca* G-kinase was present only in the soluble fraction of homogenates, as has been shown for mammalian G-kinase (Lincoln and Corbin, 1983). Phosphorylation reactions carried out on the supernatant resulted in phosphorylation of EGP-A only, but the amount of label incorporated into it was less than when the homogenate was phosphorylated before centrifugation. This reduction in labeling could be due to the existence of some factor in the 100,000 × g pellet fraction that enhanced the phosphorylation reaction. Alternatively, it could be caused by the fact that the phospho-EGP-A is more soluble than the dephospho-EGP-A.

To demonstrate that the EGPs were in fact present in the pellet fraction, we rehomogenized the pellet with the 100,000 × g supernatant taken from CNS homogenates from −24 hr animals (the EGPs are presumed absent at this time). This mixture was then phosphorylated in the presence of 0.1 mM cGMP and again separated by 100,000 × g centrifugation for 1 hr. Fluorograms of both fractions revealed the presence of both EGP-A and EGP-B in the 100,000 × g pellet and also very low levels of EGP-A in the supernatant fraction. As EGP-A was not seen when the supernatant fraction from −24 hr nervous tissue was phosphorylated, we assume that the low levels that we subsequently detected arose from a partial solubilization of the labeled EGP-A from the particulate fraction of the −4 hr nervous tissue. In the converse experiment, the pellet fraction from −24 hr nervous tissue homogenates was rehomogenized with the supernatant fraction from −4 hr nervous tissue homogenates, phosphorylated, and separated again by ultracentrifugation. Neither of the EGPs was then found in the pellet fraction and only EGP-A was found in the supernatant.

We assume that the presence of the EGPs in the particulate fraction indicates that they are somehow associated with the membrane. To see how tightly they were attached to the membrane, we tried to remove the EGPs from the pellet fraction using a number of different treatments. When NaCl was added

animals (156 ± 19%, as compared to control-injected animals; mean ± SEM, n = 8).

**Spatial distribution of the EGPs in the prepupal and pharate adult CNS**

The distribution of the EGPs in various parts of the prepupal (4 hr before ecidyse) CNS is shown in Table 1. EH stimulated an elevation of cGMP in the 3 regions of the CNS that were examined. Likewise, these 3 regions also contained the EGPs.

As with the abdominal nervous system, the brains and thoracic ganglia removed from ecidyse pupae showed no incorporation of label into the EGPs. This suggests that the EGPs in these regions of the CNS are also phosphorylated endogenously at the time of ecidyse. Also, the presence of the EGPs is not restricted to cell somata, as they were found in both the abdominal ganglia and the abdominal connectives when these were separated by dissection and homogenized and incubated separately (results not shown).

In the pharate adult, the EGPs were clearly seen in the 100,000 × g supernatant fraction of the abdominal nervous system and also in the fused pterothoracic ganglion (total homogenates). In pterothoracic ganglia removed from moths that had just ecidyse, as in the abdominal nervous system, the EGPs no longer accepted labeled phosphate. The EGPs were also seen in the pterothoracic ganglion 24 hr before ecidyse, when the animals were not behaviorally sensitive to EH. We found no definitive evidence for the presence of the EGPs in the pterothoracic ganglion or the brain. This inability was partly due to the presence of a 57 kDa protein, similar to the one found in the abdominal nervous system, which may have obscured low levels of labeling in the 54 kDa region. In some fluorograms, very faint labeling was seen in the 54 kDa region, but we could not be confident as to whether this was due to low levels of the EGPs or to smearing of the 57 kDa protein.

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Table 1. Distribution of the EGPs and cAMP-stimulated cGMP levels in the prepupal and pharate adult CNS of Manduca

| Tissue               | EGPs | cGMP (%) | Tissue               | EGPs | cGMP (%) |
|----------------------|------|----------|----------------------|------|----------|
| Brain                | +    | 174 ± 12 | Brain                | -    | 272 ± 33 |
| Thoracic ganglia     | +    | 143 ± 13 | Prothoracic ganglion (T1) | -    | 181 ± 19 |
| Abdominal ganglia    | +    | 218 ± 16* | Pterothoracic ganglion (T2-A2) | +    | 151 ± 11 |
|                      |      |          | Abdominal ganglia (A3-A7/8) | +    | 171 ± 19* |

The presence or absence of the EGPs is indicated by + or -. The increase in cGMP levels, measured 15 min after injection, in various parts of the CNS is expressed as the percentage of EH injected, compared to control animals (taken as 100% mean ± SEM of 4-8 determinations).

* Data taken from Morton and Truman (1985).

(final concentration, 0.5 M) to the homogenate after the phosphorylation reaction, and ultracentrifugation carried out, a small proportion of EGP-B was solubilized, but no effect was apparent on the distribution of EGP-A. The addition of detergents such as Triton X-100 (final concentration, 0.15%) after the phosphorylation reaction also solubilized some of the EGP-B and increased the proportion of EGP-A in the supernatant fraction (Fig. 7, C, D).

The EGPs are not the regulatory subunit of the type II cAMP-dependent protein kinase

The regulatory subunit of type II cAMP-dependent protein kinase (A-kinase) has a molecular weight of approximately 54 kDa and is phosphorylated in the presence of cAMP (Beavo and Mumbry, 1982). In bovine brain (Lohmann et al., 1980), thalamic salivary gland (McSwain et al., 1985), and Drosophila heads (Hesse and Marne, 1985), a major cAMP-dependent phosphoprotein with a molecular weight between 57 and 58 kDa was also shown to be a cAMP-binding protein by the incorporation of the photoaffinity ligand 8-azido-cAMP. Both of these properties are consistent with the phosphoproteins' being the regulatory subunit of type II A-kinase.

To determine whether either one of the EGPs was the regulatory subunit of the type II A-kinase, photoaffinity labeling was carried out with 8-azido-cAMP (Fig. 8). In prepupae, 4 hr before ecdysis, one major protein, which specifically bound 8P-8-azido-cAMP, was present in abdominal CNS homogenates. This protein was quite distinct from the EGPs, both in molecular weight (47 kDa compared to 54 kDa) and isoelectric point (4.8 compared to 5.35-5.85 and 6.45-6.75). A similar cAMP-binding protein was present in homogenates from prepupal and adult brain and abdominal nervous tissue (results not shown).

Discussion

Specificity of phosphorylation of the EGPs for cGMP

In the 3 systems studied so far, adult eclosion of Hyalophora cecropia (Truman et al., 1979), intersegmental muscle degeneration in Antheraea polyphemus (Schwartz and Truman, 1984), and pupal ecdysis in Manduca (Morton and Truman, 1985), the action of EH is mediated through cGMP. Therefore, it would be expected that the phosphoproteins involved in the action of EH would also show a clear specificity for cGMP as compared to cAMP. In homogenates of Manduca CNS, this is not the case for the EGPs. cAMP was 180 times more effective at stimulating their phosphorylation than cGMP. In Manduca CNS, the activity of the cAMP-dependent kinase is almost 10 times higher than that of the cGMP-dependent kinase (Morton and Truman, 1986), and thus may be responsible for phosphorylating the EGPs in the presence of cAMP. More comparable levels of A- and G-kinase may be found in subcompartments of the CNS, such as EH target cells.

In intact tissue, in contrast to homogenates, back-phosphorylation studies showed that cGMP was about 40 times more effective than cAMP at stimulating the phosphorylation of the EGPs. This relationship corresponds well with the finding that cGMP is more effective than cAMP at mimicking EH action (Morton and Truman, 1985). The fact that EH stimulates the phosphorylation of the EGPs, and that EH elevates cGMP and not cAMP levels in the CNS (Morton and Truman, 1985), argues that under in vivo conditions cGMP is the second messenger interposed between the EH receptor and the phosphorylation of the EGPs.

A further difference in the phosphorylation of the EGPs between homogenates and the intact CNS was the time course of phosphorylation. In homogenates, substantial levels of labeled phosphate were incorporated into the EGPs after only 5 min, whereas in the intact CNS, exposure to moderate levels of EH did not result in significant incorporation of phosphate into the EGPs until after 45 min. This delay was seen despite the fact that cGMP levels are elevated within 5 min after exposure to EH and remain elevated until after ecdysis (Morton and Truman, 1985). The time course of EGP phosphorylation in the intact CNS corresponds well with the time course of the physiological action of EH; injected EH usually takes longer than 50 min to initiate ecdysis (Truman et al., 1980). These relative times suggest that the phosphorylation of the EGPs is one of the final steps in triggering the physiological actions of EH in the target cells. The mechanism that results in such a long latency, however, is unknown.

Distribution of the EGPs throughout the CNS

The EGPs appear to be specifically phosphorylated by EH, so it would be expected that they would be located only in cells that are target cells for EH. The results presented in Table 1 show that these target cells should be found in all parts of the prepupal CNS and in the abdominal and prothoracic ganglia of the pharate adult. There were no detectable EGPs in the brain and prothoracic ganglia of the pharate adult, but this could be due to the EGPs' being obscured by other phosphoproteins, rather than to their absence. The ability of EH to elevate the levels of cGMP in these parts of the CNS suggests the presence of EH target cells.

In the above discussion we have assumed that the 2 EGPs behave as a single protein. They have the same molecular weight,
Figure 7. Fluorograms showing the distribution of the EGPs between particulate and soluble fractions of CNS homogenates. Abdominal nervous tissue was removed from animals 4 hr before pupal ecdysis, homogenized, phosphorylated with $^{32}$P-ATP in the presence of 0.1 mM cGMP, and the reaction stopped by placing it on ice. The homogenate was then centrifuged at 100,000 × g for 1 hr at 2°C, and both supernatant and pellet fractions separated by 2D SDS-PAGE, followed by fluorography. In a parallel experiment, Triton X-100 was added to the homogenate after phosphorylation (final concentration, 0.15%), vortexed, and left on ice for 1 hr. The homogenate was then centrifuged and separated as before. A, Supernatant, 100,000 × g and (B) 100,000 × g pellet fractions from phosphorylated homogenates of Manduca CNS. C, Supernatant, 100,000 × g, and (D) 100,000 × g pellet fractions from phosphorylated homogenates of Manduca CNS in the presence of 0.15% Triton X-100.

Temporal distribution of the EGPs during development
The EGPs are present before larval, pupal, and adult ecdyses. In the case of larval and pupal ecdyses, their ability to accept labeled phosphate in vitro coincides with the onset of behavioral responsiveness to EH. This is 6 hr before the normal time of larval ec dysis and 8 hr before pupal ec dysis. This close temporal correlation suggests that the lack of responsiveness to EH in appear at the same time during development, have the same spatial distribution throughout the CNS, and have the same cyclic nucleotide specificity for phosphorylation. However, they have different subcellular distributions: EGP-A is located in both soluble and particulate fractions and EGP-B is located solely in the particulate fraction of homogenates. This differential distribution could be reflected in different functions for each of the EGPs. It is also possible, however, that different posttranslational modifications of the same protein might produce different charged forms, thereby conferring on the proteins different binding properties to membranes, though the 2 proteins would retain similar functions.
Figure 8. cAMP-binding proteins in *Manduca* CNS. Abdominal nervous systems from −4 hr prepupae were homogenized and incubated with ³²P-8-azido-cAMP in the absence (A) or presence (B) of excess unlabeled cAMP. The proteins were separated by 2D SDS-PAGE and fluorograms made. The major cAMP-binding protein is marked with an arrow. C. The EGP’s do not comigrate with the cAMP-binding proteins. Abdominal nervous systems were homogenized, incubated with ³²P-ATP in the presence of 0.1 mM cGMP, separated by 2D SDS-PAGE and fluorograms made. The 2 EGP’s are marked with arrows.
each case is due to the inability to phosphorylate these proteins. Twenty-four hours before pupal ecdysis, the absence of the EGPs on fluorograms could be caused by the absence of the EGPs themselves or by their inability to be phosphorylated. We favor the former hypothesis for 2 reasons. First, the experiments that described swapping the 100,000 x g supernatants and pellets from CNS homogenates of animals 24 and 4 hr before ecdysis show that although the phosphorylating enzymes are present 24 hr before ecdysis, the EGPs are still not phosphorylated. Second, experiments with the protein synthesis inhibitor, cycloheximide, indicate that protein synthesis is necessary for the subsequent appearance of the EGPs on fluorograms. The EGPs are first visible on fluorograms 8 hr before ecdysis. Injections of cycloheximide 10 hr before ecdysis blocked the incorporation of label into the EGPs that were examined 8 hr later. Injection of cycloheximide 6 hr before ecdysis, however, did not prevent incorporation of label into the EGPs (D. B. Morton and J. W. Truman, unpublished observations). The simplest hypothesis regarding these observations is that the EGPs are absent prior to EH sensitivity in the larvae and the pupae. Their de novo synthesis is then initiated at 8–10 hr before ecdysis.

In developing adults, the EGPs are first seen 6 d before the insect is able to respond physiologically to EH. This raises the question of which step regulates EH sensitivity at adult ecdysis. All known steps in the biochemical cascade of events underlying EH action (elevation of cGMP levels and phosphorylation of the EGPs) can occur at least 20 hr before EH triggers ecdysis. Thus, at larval and pupal ecdysis the EGPs appear to be both necessary and sufficient for EH action, whereas in the adult they are necessary but not sufficient. The additional factor(s) required for adult ecdysis is (are) not known.

At each stage the ability of the EGPs to accept labeled phosphate under in vitro conditions disappears immediately at ecdysis. As with times prior to pupal ecdysis, this could be due to the absence of the EGPs or to their inability to be phosphorylated. This disappearance of the ability to label the EGPs can be prematurely induced by triggering premature ecdysis with EH or cGMP injection. We suggest that the inability to label the EGPs immediately after each ecdysis is caused by the fact that the EGPs are phosphorylated with endogenous, unlabeled phosphate as a result of the action of endogenous EH in triggering ecdysis. It is not clear what happens to the EGPs after they have been phosphorylated in vivo. In larval and pupal nervous systems the EGPs cannot be phosphorylated for several days after ecdysis, whereas in the adult, the EGPs can be phosphorylated in vitro by 24 hr after ecdysis. Because our evidence suggests that the EGPs are synthesized prior to each ecdysis, we suggest that at some point after larval and pupal ecdysis, the EGPs are degraded. This does not appear to happen after adult ecdysis. The reason for the EGPs’ continued presence in the adult CNS is unknown.

The presence of the EGPs in the very different nervous systems of the larvae, the pupae, and the adult shows that these proteins are not stage-specific. Kather, their appearance is related only to the attainment by the CNS of a specific physiological state, that of being responsive to the neuropeptide EH. As far as we are aware, this is the first example of a regulatory protein that is correlated with a particular type of behavior.

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