Sexually Dimorphic Polypeptides in Developing Antennal Sensory Neurons of an Insect

Timothy G. Kingan* and John G. Hildebrand
Arizona Research Laboratories, Division of Neurobiology, University of Arizona, Tucson, Arizona 85721

Many insects have sexually dimorphic antennae with sensilla peculiar to one sex specialized to detect sex pheromones released by the other sex. In the moth Manduca sexta, the axons of receptor cells in the male-specific sensilla as well as those common to both sexes terminate in the antennal lobes of the brain. We have used 2-dimensional electrophoresis (2DE) to study some of the proteins that are produced by sensory receptor cells in developing antennae and transported through their axons to the brain.

Extracts of antennal nerve from mature male and female M. sexta yield nearly identical 2DE patterns of proteins after staining or fluorography. Gels prepared from antennal nerves of developing animals, however, exhibit a sexual dimorphism in the quantitative pattern of at least 2 classes of proteins. One class consists of four 49 kDa proteins of similar charge, designated 49a, 49b, 49c, and 49d, and in order of decreasing mobility on nonequilibrium pH gradient electrophoresis gels. The total amount and apparent rate of synthesis of 49b and 49d are ca. 4- to 7-fold greater in antennal nerves of developing males than in those of females. Protein 49c is comparably enriched in female antennal nerves, while 49a is enriched ca. 1.2-fold in females. The second class consists of a single polypeptide of 24 kDa, which is nearly undetectable in silver-stained gels but was shown to be ca. 9-fold enriched in males by fluorography.

At the end of adult development, male and female moths have similar patterns of the 49 kDa polypeptides, and synthesis of the 24 kDa polypeptide is reduced to nearly undetectable levels. The patterns of sexual dimorphism thus appear to be associated with the growth and maturation of antennal sensory axons into the antennal lobes. Biosynthesis of the 24 kDa polypeptide was nearly undetectable in antennal nerves or fragments of antennal nerves incubated in vitro with radiolabeled methionine. The 49 and 24 kDa polypeptides appear to belong to separate classes of rapidly transported proteins. The 24 kDa polypeptide is among the most rapidly transported proteins; it is found exclusively in a particulate fraction and is associated with plasma membrane but apparently not mitochondria. The 49 kDa polypeptides are found in both the particulate and soluble fractions; the more basic 49a and 49b are enriched in the particulate fraction, while the more acidic 49c and 49d are enriched in the soluble fraction.

Gender-specific behaviors and reproductive functions of many animals have anatomical correlates in their CNS. Such sexual dimorphism, which can be manifested as differences in cell number, size, or extent of arborization in certain brain nuclei, underlies sex-specific behavior in the adult (Arnold and Gorski, 1984). In vertebrates, gonadal steroids act perinatally and neonatally to control the development of sexually dimorphic structures in the brain; much earlier the same steroids regulate the dimorphic development of the reproductive system. Thus, while the gonads themselves are genetically determined, all other sexual dimorphism depends on gonadal secretions.

In many invertebrates sex-specific behaviors have been well documented, but relatively little is known about their cellular or developmental bases in the nervous system. For example, many male moths exhibit species-specific, stereotyped behaviors when exposed to the sex pheromones released by conspecific females. The male moths respond with characteristically patterned, upwind flight in the plume of pheromone in search of a female. The pheromone-detecting sensory cells innervate sensilla in the antenna. Often, as in the sphinx moth Manduca sexta, the antennae are sexually dimorphic, and the pheromone-detecting receptor cells are associated with long, male-specific olfactory hairs on the antenna, the sensilla trichodea (Sanes and Hildebrand, 1976a). The olfactory receptors of antennal sensilla send out axons, up to 2 cm long, that project through the antennal nerve (AN) and terminate in the antennal lobe (AL) of the brain (for review, see Hildebrand, 1985). It has been shown in M. sexta that axons from the trichoid sensilla terminate exclusively in a male-specific synaptic region of the AL, the magroglomerular complex (MGC).

The antennae of moths arise during metamorphic adult development from imaginal disks present in the head of the larva. The sexual dimorphism of the antenna is genetically determined, so that imaginal disks can be transplanted between sexes to give rise to antennae that are typical of the donor's gender (Schneiderman et al., 1982; Hildebrand, 1985). The structure of the AL, however, is not genetically determined: The MGC characteristic of the normal male develops only in an AL innervated by sensory axons from a male antenna. Indeed, if a female receives a male antennal disk by transplantation, she

* Present address: Peptide Design, 13331 Middlebrook Road, Germantown, MD 20874.
develops an MGC in the AL innervated by the AN from the grafted antenna (Schneiderman et al., 1982; Hildebrand, 1985). Such transplantation experiments have revealed that development of the MGC depends upon direct innervation of the AL by male-specific antennal axons. Thus, the MGC does not develop if the male AN fails to contact the brain or does so at sites other than the AL. This requirement for intimate interaction between the AN and the AL raises the possibility that one or more nondiffusible substances, synthesized in male antennal sensory neurons and transported along their axons, may influence the formation of the MGC. Likely candidates for this role would be polypeptides unique to, or highly enriched in, male ANs. We have pursued this possibility by comparing the total and axonally transported polypeptides in ANs of developing male and female M. sexta by 2-dimensional electrophoresis (2DE). A preliminary account of some of these results appeared previously (Kingan and Hildebrand, 1983).

**Materials and Methods**

**Animals**

*Manduca sexta* (Lepidoptera: Sphingidae) were reared at 26°C and ca. 60% relative humidity on artificial diet (modified from that of Bell and Joachim, 1976) in a long-day photoperiod (17 hr light:7 hr dark). Insects were staged according to previously published criteria (Sanes and Hildebrand, 1976a; Tolbert et al., 1983). Adult development occurs over 18–21 d, corresponding to ca. 21 d.

**Radiolabeling of AN proteins**

Proteins in the ANs of developing moths were labeled with L-35S-methionine by in situ and in vitro procedures.

**In situ labeling.** L-35S-Methionine (480–2000 Ci/mmol, Du Pont, NEN Research Products) was injected into male pupae reared at 26°C in a glass microhomogenizer in preparation for extraction and electrophoresis.

**In vitro labeling.** Radiolabeled methionine was dried as described above and redissolved in sterile methionine-free Grace's medium.Brains with attached ANs were removed from developing animals and placed in a drop of methionine-free Grace's medium for 15 min. They were then transferred to 50 μl Grace's medium containing L-35S-methionine (50 μCi, >1000 Ci/mmol) and incubated in a hanging drop for 5 hr. The brains were then transferred to medium without radiochemical and rinsed for 15 min. The AN was then separated from the AL and placed in a glass microhomogenizer in preparation for extraction and electrophoresis.

**Subcellular fractionation**

Two procedures were used for tissue fractionation. For preparation of total particulate and soluble fractions, ANs and ALs were disrupted in a glass microhomogenizer at 0°C in buffered medium containing protease inhibitors of the following composition (modified from Shorr et al., 1981): 10 mM Tris-HCl, 2.5 mM EDTA, 2.5 mM EGTA, 0.1 mM benzamidin chloride, 1 mM benzamidine, 100 μg/ml Bactristatin, 5 μg/ml soybean trypsin inhibitor, and 2 μg/ml leupeptin, pH 7.4 (all reagents obtained from Sigma Chemical Co.). The homogenate was separated into soluble and particulate fractions by centrifugation at 4°C for 5 min. The supernatant (soluble) fraction was decanted and held on ice, and the pellet (particulate fraction) was dried by lyophilization. To precipitate proteins in the soluble fraction, cold (-20°C acetone:vol) was added with mixing. The resulting precipitate was isolated by centrifugation, washed once with 80% acetone, and dried briefly in the vacuum centrifuge. The dried particulate and soluble fractions were stored at -80°C no longer than 2 hr prior to analysis.

For fractionation of total membranes, whole brains or brains lacking optic lobes and subesophageal ganglia (in either case, with ANs and ALs still attached) were disrupted in a Potter-Elvejhem homogenizer in 5 mM Tris-1/2 mM sucrose, pH 7.4, with 30 strokes of the rotating Teflon pestle (Polyscience Corp. homogenizer, model PSC10, Invert setting) in 1 min at 0°C. The homogenate was centrifuged at 800 × g for 10 min (Sorvall SS-34 rotor), and the pellet was resuspended in Tris/sucrose and centrifuged as above. The supernatant fractions were pooled and ANs and ALs pelleted as above for total particulate material (Beckman SW 50.1 rotor, 35,000 rpm, 60 min). The supernatant fluid was removed and retained for analysis. The pellet was resuspended in a small volume of 5 mM Tris-1/2 mM sucrose, pH 7.4, and layered onto a discontinuous sucrose gradient containing layers of 20, 26, 31, 37, 45, and 55% sucrose (Lorenz and Willard, 19/8) or 20, 31, and 55% sucrose in 5 mM Tris-HCl, pH 7.4. Membranes were then centrifuged for 7.8 × 105 g, min (Beckman SW 41 ti, 26,500 rpm, 90 min). The bottom of the tube was punctured, and the interfaces were collected directly into ultracentrifuge tubes. The contents were diluted with ice-cold 5 mM Tris and 5 mM Tris-1/2 mM sucrose with mixing, and the membranes were pelleted as above for total particulate material. Membranes were resuspended in 50 mM Tris-HCl, pH 7.4, for enzyme assays or in water for lyophilization and subsequent electrophoresis.

**Enzyme assays**

Standard procedures were used to quantify ouabain-sensitive Na-K ATPase (EC 3.1.3.1, Medzhitov et al., 1971; Breer and Knipper, 1985), succinate dehydrogenase (EC 1.3.99.1, Porteous and Clark, 1965; Breer and Knipper, 1985), and protein (Bradford, 1976).

**Electrophoresis**

2DE was performed using non-equilibrium pH gradient electrophoresis (NEPHGE; O'Farrell et al., 1977). Tissue was solubilized directly in lysis buffer (O'Farrell et al., 1977) and then centrifuged at 8000 × g for 5 min. When samples were to be compared within an experiment, equal amounts of radioactivity were loaded on gels. Radioactivity was determined by adding 1 μl aliquots of samples to 200 μl of 0.1% BSA; the mixture was precipitated with 8% trichloroacetic acid. After rinsing and drying, precipitates were solubilized in Protosol (DuPont, NEN Products) prior to addition of liquid scintillation cocktail.
programs to obtain information about the relative accumulation of 35S-methionine in individual polypeptides.

**Results**

Initial experiments using SDS-PAGE alone or with isoelectric focusing (O'Farrell, 1975) in 2DE failed to reveal sexually dimorphic proteins in extracts of ANs of developing moths. Because at least some basic proteins did not enter the gels, we repeated the experiments using NEPHGE-2DE. This procedure allowed us to detect a greater number of polypeptides in silver-stained gels; in addition, when comparing extracts of male and female ANs, we found significant quantitative differences among certain polypeptides.

The patterns of stained proteins obtained with extracts of combined ANs and ALs from developing males and females are nearly identical (Fig. 1A). Nevertheless, we could reproducibly detect a group of 4 polypeptides, all with apparent molecular weights of 49 kDa and apparently similar charge, that differ quantitatively in male and female ANs and ALs (Fig. 1B, C). These polypeptides are designated 49a, 49b, 49c, and 49d in order of increasing acidity (left to right in the photographs). As can be seen in Figure 1B, polypeptides 49b and 49d are enriched in male ANs and ALs, while 49a and 49c are enriched in female ANs and ALs. These differences in the stainable polypeptides can be detected at least as early as stage 5/6, the youngest animals used in these experiments. When extracts of ANs and ALs were electrophoresed separately, we did not detect significant differences in the relative amounts of these polypeptides.

Axons in the AN are bundled in fascicles surrounded by glial-like cells (Sanes and Hildebrand, 1976b). These ensheathing cells contribute proteins to the stained pattern and could conceivably account for some of the observed sexual dimorphism.

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**Figure 1.** NEPHGE-2DE (this and all other gels) of AN and AL tissue from stage 11 developing adults; proteins visualized by silver staining. A, left: male; right: female; NEPHGE gels were run side-by-side on an SDS gel. B (male) and C (female). Details of appropriate 49 kDa regions from A shown in brackets. The female sample contained ca. 50% more protein than that in the male sample. The 49 kDa polypeptides are designated 49a, 49b, 49c, and 49d in order of decreasing mobility on NEPHGE, i.e., apparently increasing acidity. The positions of molecular-weight markers (from a separate gel) are indicated to the right in A and are, from top to bottom (in kDa): β-galactosidase, 116; phosphorylase B, 97.4; BSA, 66; ovalbumin, 45; carbonic anhydrase, 29; lysozyme, 14.
Figure 2. Fluorogram after 2DE of extracts of in situ labeled ANs and ALs of 2 males and 2 females. 35S-Methionine was injected into the distal third of antennae of stage 9 male and female developing adults. ANs and ALs were removed 2.5 d later, at which time the animals had developed to stage 11. A, Male (left) and female (right) extracts, run side-by-side in the second-dimension (SDS) gel. Details of the 49 kDa region are shown for the male (2B) and the female (2C). Molecular-weight markers (to the right) as described for Figure 1.

To test this possibility and to sample the complement of proteins synthesized and transported at discrete stages of development, we labeled proteins by injection of 35S-methionine into the antennae of developing moths. The ipsilateral AL and the intracranial portion of the corresponding AN were removed 0.5-3 d after the injection. The results of a 2.5 d experiment are shown in Figure 2. The 49 kDa proteins are present in the fluorograms from both male and female ANs and ALs (Fig. 2 B, C); in addition, their patterns of relative abundance are qualitatively similar to those in the stained gels (cf. Fig. 1, B, C).

The fluorograms reveal an additional polypeptide, of 24 kDa apparent molecular weight, that is enriched in extracts of male ANs and ALs (Fig. 2 A). Because silver-stained gels only occasionally show a very faint spot in this position, it seems that this protein either is present in very small amounts relative to others or is refractory to staining by our methods.

Our initial findings showed that ANs contain 5 polypeptides, synthesized in developing moths and differing quantitatively in males and females. These polypeptides apparently are synthesized in sensory cell bodies in the antenna and move to the AN and AL by axonal transport. Alternatively, some of the polypeptides in the fluorograms might have been produced in the glial-like cells of the AN or cells in the AL, incorporating free 35S-methionine that had reached the AN or AL. We attempted to address this possibility by examining the complement of polypeptides synthesized by ANs and ALs from stage 7-8 males in short-term organ culture (see Materials and Methods). While many proteins found in the "axonal transport" fluorograms were also found in the "organ culture" fluorograms, the overall patterns were very different (cf. Fig. 3, A, B), supporting our conclusion that injections into the antenna labeled only transported proteins. By overlapping fluorograms we found spots in the "organ culture" fluorograms corresponding to all 49 kDa polypeptides, although their relative labeling seemed not to correspond to either the male or the female pattern. In addition, we found a spot corresponding to the 24 kDa polypeptide, but its intensity in "organ-culture" fluorograms was very low, particularly with respect to the total level of protein biosynthesis in...
ANs in vitro (Fig. 3, B, C). Similar fluorograms prepared from AL extracts also reveal synthesis of the 49 kDa polypeptides; we could not, however, detect the 24 kDa polypeptide in these preparations (Fig. 3D). Thus, the in vitro studies show that the relative amounts of the 49 kDa polypeptides reaching the AL by axonal transport are markedly greater than those arising by local synthesis. Moreover, the 24 kDa polypeptide may be completely absent (AL) or present in only minute amounts (AN) in preparations monitoring local biosynthesis. We conclude that the 49 and 24 kDa polypeptides revealed after in vitro labeling were proteins synthesized in antennal sensory neurons and transported through their axons in the AN toward their terminals in the AL.

Knowledge of the rates of transport of axonal polypeptides is helpful if inferences about their function are to be made. We therefore sought to determine the relative rates of transport of the 49 and 24 kDa polypeptides by removing the ANs and ALs from animals at intervals of 0.5–3 d after the injections. Unfortunately, this antennal preparation does not lend itself to absolute determinations of rates of axonal transport. Somata of antennal receptor cells are distributed along the entire length of the antennal flagellum. If radiolabeled amino acid injected focally into one annulus of the flagellum spreads to several other annuli, resolution of different transport classes of polypeptides is correspondingly decreased. Nevertheless, when the ANs and ALs are removed from animals 0.5 or 1 d after the injection, far fewer radiolabeled polypeptides are found than in the 2 or 3 d preparations (Fig. 4). Apparently, the more rapidly transported polypeptides account for the radiolabel in the 0.5 or 1 d extracts. Among these relatively rapidly transported species is the 24 kDa polypeptide, which represents a greater proportion of the total label in the 0.5 or 1 d extracts than in 2 or 3 d extracts (Table 1). Of the 49 kDa polypeptides, only 49a can be visualized at 24 or 36 hr after injection. The other 49 kDa polypeptides appear in the ANs and ALs at 48 hr, however, along with many other polypeptides not previously detected.
Thus, the 24 kDa polypeptide belongs to a class of rapidly transported polypeptides, and the bulk of the 49 kDa polypeptides belongs to a different, more slowly transported class.

An estimate of the relative rates of synthesis of the 24 and 49 kDa polypeptides in males and females was obtained after scanning the fluorograms from extracts prepared 2.5 d postinjection (Fig. 5). The 49b and 49d polypeptides are enriched ca. 3.5- and 6.7-fold, respectively, in males, while 49a and 49c are enriched ca. 1.2- and 2.4-fold in females. The enrichment of 49a is ca. 4-fold when total protein is examined with silver-stained gels (data not shown). Small enrichments in the synthesis and transport of specific polypeptides may be amplified by factors controlling turnover of protein in axons and nerve terminals.

Table 1. Appearance of label in 24 kDa polypeptide following injection of 35S-methionine

| Hours post injection | Percentage 24 kDa polypeptide* |
|----------------------|-------------------------------|
| 24                   | 1.73 ± 0.10*                  |
| 48                   | 0.99                          |
| 63                   | 0.45                          |
| 48 (stage 16)        | 0.15                          |

* Value reported is the percentage of the total optical density in the gel represented by the 24 kDa polypeptide. Animals were stage 7/8 at time of injection, except as indicated.

If the 24 and 49 kDa polypeptides are to be considered candidates for a role in development, it is important to determine their distributions among subcellular fractions. Accordingly, we prepared total particulate and soluble fractions from ANs and ALs for 2DE (see Materials and Methods). All 49 kDa polypeptides are found in the particulate fractions (Fig. 8A). The soluble fraction (Fig. 8B) is greatly enriched in 49c and 49d,
The 24 and 49 kDa polypeptides were quantified from X-ray film after fluorography, and normalized ratios (see Materials and Methods) are reported. Data are from experiment reported in Figure 2.

while 49a and 49b are almost undetectable. The 24 kDa polypeptide is found in the particulate fraction but not in the soluble fraction.

We have extended these findings by analyzing the total-membrane fraction from ANs and ALs on sucrose gradients. In a preliminary study designed to determine the distribution of marker enzymes, we fractionated total membranes from the brains of stage 7/8 males on discontinuous gradients consisting of 20, 26, 31, 37, 45, and 55% sucrose (see Materials and Methods). Ouabain-inhibitable Na/K-dependent ATPase activity (ATPase) is enriched at the surfaces of 26 and 31% sucrose, while most of the succinate dehydrogenase activity (SDH) passes through 37 and 45% sucrose and is enriched at the surface of 55% sucrose. On the basis of these findings, we chose to fractionate the less abundant tissue from animals receiving injections of 35S-methionine with step gradients containing 20, 37, and 55% sucrose. The distributions of ATPase, SDH, and the 24 kDa polypeptide are shown in Figure 9. The enrichment of SDH in the “mitochondrial” fraction (37/55) is ca. 7-fold, while the ouabain-sensitive ATPase is only slightly enriched in the lighter membrane fraction (20/37). Nevertheless, the 24 kDa polypeptide is ca. 10-fold enriched in the 20/37 fraction; this fraction is likely to include plasma and/or microsomal membranes. The 24 kDa polypeptide is not associated with the heavier, “mitochondrial” fraction.

Discussion
The antennal flagellum of both male and female M. sexta comprises about 80 annuli. While the 2 basal segments of the antenna (scape and pedicel) contain numerous mechanosensory sensilla, the sensilla in the flagellum are believed to be predominantly chemosensory. Approximately 56,000 sensilla basiconica in male M. sexta (Sanes and Hildebrand, 1976a) closely resemble the most abundant sensilla in females. Sensory neurons innervating sensilla of this class respond to plant-derived volatile substances. In addition, each antenna of male M. sexta contains ca. 43,000 sensilla trichodea (Sanes and Hildebrand, 1976a) whose sensory cells respond to female pheromones (Kassling et al., 1989). These receptor cells contribute 25–33% of the axons in the AN of the male. Thus, although the AN contains axons of several different functions, comparison of the proteins in male and female ANs should reveal candidates for male-specific functions.

The patterns of polypeptides extracted from ANs and ALs of developing males and females and visualized after 2DE were nearly identical. Using NEPHGE-2DE, we identified 2 classes of polypeptides, distinguished on the basis of apparent molecular weight, in which differences between males and females are reproducibly demonstrated. One class contains four 49 kDa polypeptides of similar charge, and the second consists of a single 24 kDa polypeptide.

These 2 classes of polypeptides also differ in apparent rates of axonal transport. The patterns of labeling of different groups of polypeptides extracted from ANs on successive days following injection of 35S-methionine provide evidence for at least 2 classes of rapidly transported proteins. Newly synthesized 24
kDa polypeptide appears in the AN 12 hr after injection of \( ^{35} \text{S}- \) methionine into the antenna 20 mm distal to the intracranial portion of the AN analyzed in these studies. At this time, the majority of the labeled proteins are found as a series of spots in arcs at the top of the gel, suggestive of glycoproteins. Among the 49 kDa polypeptides, a small amount of 49a was detected in the 24 hr fluorograms, but the bulk arrives at 48 hr along with the other 49 kDa polypeptides. They arrive in the AN at the same time as many other labeled proteins (cf. Figs. 2 and 4). The majority of these proteins arrives 36-48 hr after injection, perhaps as a wave-front comprising a distinct class of transported proteins.

The sexually dimorphic patterns of the 24 and 49 kDa polypeptides are most pronounced during early development, beginning no later than stages 5/6 of adult development and extending at least until stage 13. Sensory axons begin to reach the AL at stage 3 and have begun to grow into the AL neuropil by stage 4 (Camazine and Hildebrand, 1979). Formation of AL glomeruli (including the MGC) proceeds from stage 4 and is complete by stage 13 (Tolbert et al., 1983; Schneiderman, 1984). Thus, elevated synthesis of polypeptides 24 kDa and 49b and 49d in males is demonstrable at a time in development coincident with the growth of sensory axons and their participation in formation of glomerular synaptic neuropil. The sexual dimorphism in the expression of the 24 kDa polypeptide ends late in development; by stage 16, its synthesis and transport are reduced to nearly undetectable levels. In this respect, the 24 kDa polypeptide resembles the growth-associated proteins demonstrated in axons arising from toad and rabbit retinal ganglion cells (Skene and Willard, 1981a, b), one of which was reported to be either 24 kDa (toad) or 23 kDa (rabbit) in molecular weight. The 24 kDa polypeptide described in this report appears to be synthesized only at very low levels or not at all by CNS tissue (AL) or fragments of AN containing glia but not the somata of sensory cells. Of the 49 kDa polypeptides in *M. sexta*, 49d shows a significant decline in incorporation of radioisotope late in development in males; 49a, 49b, and 49c are still readily detectable when labeling is carried out 2 d before emergence of the adult moth. The stained gels reveal much less sexual dimorphism in these polypeptides than is demonstrable early in development.

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**Figure 8.** Particulate (A) and soluble (B) fractions were subjected to 2DE and fluorography. The 24 kDa polypeptide is restricted to the particulate fraction; there is a slight "tailing" of a polypeptide running immediately below the expected location of the 24 kDa polypeptide. The 49 kDa polypeptides are present in both the particulate and soluble fractions. 49a and 49b (black arrowheads) are enriched in the particulate fraction; 49c and 49d (white arrowheads) are enriched in the soluble fraction.

**Figure 9.** Membranes were collected at the interfaces of 20-37% (20/37) and 37-55% (37/55) sucrose. Aliquots were assayed for SDH and ouabain-inhibitable ATPase activities, while additional aliquots were subjected to 2DE and fluorography for quantification of the 24 kDa polypeptide. P<sub>i</sub>, phosphate.
development. The patterns in the stained gels suggest that females accumulate 49b and 49d late in development, reducing the sexual dimorphism in the staining patterns.

The sexually dimorphic polypeptides discussed in this report are apparently different from the male-specific antigen detected in the sensory cells of the trichoid sensilla of *M. sexta* with a monoclonal antibody (Hishinuma et al., 1988a). This antigen, termed male olfactory specific antigen (MOSA), is detectable in male sensory cells only late in development (Hishinuma et al., 1988b). The fact that we have not detected a protein in the gel patterns from ANs and ALs of mature moths that is a candidate to be MOSA may be due to limitations of 2DE. It is also possible that modifications by male-specific sensory cells of a protein also present in non-sexually-dimorphic cells alters its immunogenicity to produce MOSA without appreciably changing its electrophoretic properties.

A number of polypeptides not discussed in this report decline in synthesis and/or transport late in development with time courses similar to that of the 24 kDa polypeptide. While we have not detected sexually dimorphic expression of these polypeptides, they are candidates for a role in the development of synaptosomal interactions in the "ordinary" glomeruli of the AL.

The firm association of the 24 kDa polypeptide with the particular fraction and the enrichment of that polypeptide in plasma membrane/microsomal fractions is consistent with its rapid rate of axonal transport. These findings support the possible involvement of the 24 kDa polypeptide in developmental events leading to the formation of the MGC. The subcellular distribution of the 49 kDa polypeptides is predicted, to some extent, by their apparent charge as revealed in NEPHGE: the more acidic 49c and 49d are greatly enriched in the soluble fractions, while 49a and 49f are found in the particulate fractions. It is not yet known if the 49 kDa polypeptides are related to each other, as suggested by the apparent identity of their molecular weights and the similarity of their charges (mobility on NEPHGE gels).

Because of their sexually dimorphic, pheromone-sensitive sensory cells, males have more afferent axons in their AN than do females. If the higher rate of synthesis of the 24 kDa polypeptide is attributable to the male-specific sensory cells, then its enrichment in these cells would be significantly greater than 9-fold, which was determined for whole ANs. The enrichment of 49b and 49d in males could be accounted for in a similar way. The enrichment of 49a in females, while ca. 4-fold on stained gels, is only 20% on the fluorograms; 49c appears to be similarly enriched in females ANs upon quantification in stained gels (2.6 x) and fluorograms (2.4 x). It is difficult to explain the different quantitative enrichments in females of 49a and 49c solely on the basis of a depletion in male ANs. It is possible that these proteins are subject to different rates of turnover.

The functions of the 49 and 24 kDa polypeptides are unknown. Identification of molecular differences in the proteins synthesized by male and female antennal receptor cells is, nevertheless, compelling in its implications for the study of neural development. Male-specific, functionally specialized neuropil structures can develop in ALs of either sex, but only when some cue from sensory afferents of the male antenna is provided at close range (Schneiderman et al., 1982). Moreover, the male AL has ca. 30 more nerve cell bodies than the female AL in the group of somata known to contain male-specific MGC projection neurons (Matsumoto and Hildebrand, 1981; Christensen and Hildebrand, 1987; Homberg et al., 1988). It seems likely that axons from trichoid sensilla influence these neurons or others common to both sexes to form the MGC. The accessibility of the developing peripheral and central elements of the moth olfactory system provides an unusual opportunity to investigate the roles of specific polypeptides in directing the development of a sexually dimorphic neuropil.

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