CELLULAR INTERACTIONS AND PATTERN FORMATION IN THE VISUAL SYSTEM OF THE BRANCHIOPOD CRUSTACEAN, DAPHNIA MAGNA

III. The Relationship Between Cell Birthdates and Cell Fates in the Optic Lamina

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Abstract

The birthdates of nerve cells in the optic ganglion of the water flea Daphnia magna were determined using [3H]thymidine autoradiography. The analysis was aided by computer-assisted three-dimensional reconstruction and correlated with serial electron microscopy of the developing visual system. Previous work has shown that the projection from eye to optic ganglion is retinotopic at the level of single cells (Macagno, E. R., V. LoPresti, and C. Levinthal (1973) Proc. Natl. Acad. Sci. U. S. A. 70: 433-437). Studies of both normal and perturbed development (LoPresti, V., E. R. Macagno, and C. Levinthal (1973) Proc. Natl. Acad. Sci. U. S. A. 70: 56-61; Macagno, E. R. (1978) Nature 275: 318-320) have suggested that retinotopy is a consequence of an invariant pattern of sequential growth and cellular interactions. We report here that the laminar cells, the first-order target cells of the visual projection, become postmitotic in an orderly fashion that correlates with their position and connectivity in the adult array. Furthermore, presumptive laminar cells complete their final division just prior to or during initial contact with the growing processes of the visual fibers with which they will form their mature retinotopic contacts. The correlation between times of final division of target cells and the arrival of visual fibers with which they interact provides further indirect evidence that sequential growth and cellular interactions are key factors in establishing normal connectivity in the visual system of arthropods.

Ordered patterns of cell proliferation are common features in the embryogenesis of neuronal structures. Well known examples of topographically organized patterns of nerve cell birthdates include the retina and optic ganglion (OG) of insects (Nordlander and Edwards, 1969b; Hofbauer and Campos-Ortega, 1976; Ready et al., 1976; Anderson, 1978; White and Kankel, 1978; Mouze, 1980; Stark and Mote, 1981), the segmental ganglia in both grasshopper (Bate, 1976) and leech (Weisblat et al., 1980), the retina and tectum of birds and amphibians (Straznicky and Gaze, 1971, 1972; LaVail and Cowan, 1971), and the cerebral and cerebellar cortices (reviewed by Sidman and Rakic, 1973) as well as the spinal motor columns in mammals (reviewed by Hollyday, 1980). The patterns of neuronal birthdates observed in these various examples vary considerably in their geometry and in other respects. The relevance of these birthdate patterns to either local or distant circuit features of the adult structure is largely unknown.

The structure and development of the projection from eye to OG in a single clone of the waterflea, Daphnia magna, has been described in detail previously (LoPresti et al., 1973; Macagno et al., 1973; Flaster et al., 1982). Briefly summarized, the visual system consists of a single, bilaterally symmetric compound eye composed of 22 fused-rhabdom ommatidia projecting via the optic nerve to an OG which contains two groups of neurons and a small number of glial cells. On the ganglion's anterior surface are the cell bodies of the laminar neurons, the first-order cells of the ganglion. More posteriorly situated...
are the cell bodies of the medullary neurons which are predominantly second- or higher-order interneurons. The eight photoreceptors of each ommatidium project their axons in a fascicle to the optic lamina where, together with five laminar cells, they form a unit structure, the optic cartridge. There are 22 optic cartridges. Each ommatidium projects exclusively to a single laminar cartridge in a retinotopic manner with the majority of photoreceptor-laminar cell synapses made between the members of a single cartridge. The structure of the visual system and the projection from compound eye to lamina are diagrammed in Figure 1.

Serial section electron microscopic studies of the developing visual system of *Daphnia* have shown that the
ommatidia mature in an orderly, lateral-to-medial sequence (see Fig. 1a, lower diagram). Within each ommatidium, the axons of the photoreceptors grow as a fascicle composed of a leading fiber with an enlarged growth cone and seven followers with smaller growth cones. Each lead fiber grows along a midline glial palisade which bridges a narrow extracellular sinus, enters the primordial lamina, and interacts sequentially with five presumptive laminar cells. The laminar cells wrap around the lead fiber and, shortly thereafter, each cell generates a neurite which joins the fascicle of optic axons. The interaction between lead fiber and laminar cell, culminating in its differentiation as a member of that cartridge, we have called recruitment. The first optic cartridges formed come to be situated laterally in the lamina; the last cartridges to form are situated medially (see Fig. 1a, lower panel). The ingrowth of the projection from eye to lamina and the formation of nascent cartridges is completed in about 10 hr (at 22°C) during the middle third of embryogenesis. Studies using a UV microbeam to delete photoreceptors or to delay the arrival of optic axons into the laminar primordium have demonstrated that embryonic laminar cells are dependent upon interactions with optic axons for their survival and differentiation (Macagno, 1979, 1981). When a few photoreceptor axons are caused to arrive a few hours late, the laminar cells with which they interact show a corresponding delay in their own development, indicating that differentiation of laminar neurons is triggered by an interaction between optic axons and presumptive laminar cells.

In this study we ask: when do presumptive laminar neurons become postmitotic? Is there an orderly arrangement of laminar cell birthdates with respect to the cartridges to which these cells come to belong? Finally, what is the temporal relation between the production of laminar cells and their interaction with incoming optic axons?

Materials and Methods

Embryos used in these experiments were all taken from a single clone of *Daphnia* cultured under standard conditions at 22.5 ± 1°C (Macagno et al., 1973). Embryonic age was established by time elapsed after the extrusion of eggs from the ovaries into the dorsal brood pouch. Embryos were removed from the brood pouch between 22 and 24 hr after extrusion. They were maintained in a culture medium containing penicillin and streptomycin (Murphy, 1970) until the onset of sustained swimming, at about 60 hr of age, at which time the animals were transferred to normal culture medium. By 60 hr, development is essentially complete.

For microinjection, embryos were transferred to physiological saline immediately before injection and were returned to Murphy's medium 4 to 6 hr later. Pressure-driven microinjections were accomplished through glass micropipettes with 2- to 2.5-μm tip diameters using a device capable of brief pressure pulses (McCaman et al., 1977). The output of the pipettes was estimated by measuring the amount of radiolabel ejected into water by test pipettes. Roughly 100 pl of solution were delivered into each embryo using pulses of 50 msec duration at a pressure of 40 psi. For birthdating, the injection solutions consisted of [methyl-6-H]thymidine, 40 to 60 Ci/mmol (New England Nuclear or Amersham) at a radiochemical concentration of 2.5 mCi/ml in *Daphnia* physiological saline. Methylene blue (final concentration 0.5%) was added to make injections visible. Embryos were held for injection by placing them in small drops of water on wet tissue paper mounted on glass slides. The micropipette was guided by means of a Leitz micromanipulator and was inserted into the dorsoposterior, predominantly yolky region of the embryos. The injections were performed under a compound microscope equipped with a rotating stage to facilitate orientation of the embryos. In birthdating experiments, each embryo received a single injection. The injection was made at one of a number of developmental stages during the middle third of embryogenesis (23 to 43 hr of development). The injected animals were fixed 4 to 5 days later, well into the juvenile instars. Some specimens were fixed shortly after the introduction of label. All animals were fixed and embedded in Epon-Araldite as previously described (Macagno et al., 1973). For autoradiographic analysis, specimens were sectioned serially at 1 μm in the case of juveniles and 0.5 μm in the case of embryos. The sections were dried on glass slides, dipped in Kodak NTB-3 emulsion (2 parts emulsion: 1 part distilled water), and exposed at 4°C for 14 to 16 days for 1-μm sections and for 25 days for 0.5-μm sections. The emulsions were developed with Kodak D-19 and fixed, using the manufacturer's recommended protocol. The sections were then stained through the emulsion with 1% toluidine blue, followed by a brief destaining in a dilute aqueous solution of acetic acid and ethanol. Serial section autoradiographs were photographed with an oil immersion lens on a Zeiss photomicroscope. The resulting negatives were successively aligned and photographed to produce filmstrips suitable for computer-assisted three-dimensional reconstruction using the CARTOS system (Macagno et al., 1979). For laminar neurons, laminar glial cells, and medullary neurons, the positions of their nuclei were recorded, and the number of silver grains over each nucleus was counted. A set of computer programs was written which would then count the number of cells traced, generate histograms of the distribution of cells as a function of normalized grain counts, and, on graphics devices, display cells in groupings determined by their label intensity (histogram classes).

Analysis of either control series (embryos which did not receive [3H]thymidine injection) or nonlabeled areas from thymidine-injected specimens showed equivalent grain densities, two to three orders of magnitude below grain densities over labeled cell bodies.

For correlated electron microscopy, embryos of the same age as those injected for autoradiographic study were fixed, stained en bloc with uranyl acetate, and embedded in Epon. These specimens were serially thin sectioned (at thicknesses showing silver to gold interference colors), and every third or fourth section was photographed at × 1,000 magnification on a Zeiss EM-9 electron microscope. The numbers of optic fiber bundles reaching the optic laminar primordium were counted directly from serial prints. Other aspects of the embryonic lamina were also examined using these same serial
The pattern of \(^3\)H thymidine labeling seen in embryos. Embryos from a single brood were injected with \(^3\)H thymidine at 34.5 hr in development, fixed after an interval of either 1.5 or 4 hr (at 36 and 38.5 hr, respectively), processed for serial section autoradiography, and studied in serially aligned prints. The number, position, and characteristics of labeled cells in and around the laminar primordium were noted. For comparison, and to visualize features not resolved with the light microscope, embryos at the same stages in development were sectioned for electron microscopy.

At 36 hr, \(1\frac{1}{2}\) hr after the \(^3\)H thymidine injection, labeled cells are seen in the posterior laminar primordium, in the medullary primordium, and in the primordium of the supraspinal ganglion. Cells in the anterior lamina that appear already recruited into cartridges are unlabeled. In one reconstructed specimen, the number of unlabeled cells is 23/hemilamina, and roughly half of them appear to be in cartridges. As will be seen later, the number of unlabeled cells is not too different from the number obtained from animals injected at 34 hr but fixed when adult. Posterior to the region of unlabeled cells, heavily and moderately labeled cells are present both on and near the midline. Most of these cells are of uniform size. A few mitotic figures can be recognized, and most of these are labeled. A mitotic figure in the midline region, where presumptive laminar cells are first recruited, can be seen in the electron micrograph of an embryo fixed at 35 hr of development as shown in Figure 2c. Larger cells, many of them labeled, make up a major part of the posterior region of the optic ganglion primordium. A laminar-medullary boundary is not readily distinguishable at this stage, precluding the identification of possibly separate precursor cells for these structures.

In a specimen fixed 4 hr after injection of label (38.5 hr in development), the number of photoreceptor fiber bundles has increased, as has the area in the lamina composed of cartridges. Two autoradiographs from a specimen fixed at 38.5 hr are shown in Figure 2, a and b. Counts of unlabeled cells show 29 cells on one side of the lamina and 28 on the other, numbers in excellent agreement with the number of unlabeled cells seen in adults injected at 34 hr of age (averaging 28 cells/hemilamina) and not very different from the number of unlabeled cells counted at 36 hr (23 cells/hemilamina). The majority of these unlabeled cells appear to be in cartridges. As one moves posteriorly through the lamina, labeled cells begin to appear but are confined to the midline, surrounded laterally by unlabeled recruited cells (Fig. 2b). Still further posteriorly, nearly all cells are labeled.

These results suggest that cells already recruited into nascent cartridges are postmitotic, whereas cells not yet withdrawn from the cell cycle are not yet recruited into cartridges and are located posteriorly and medially in the laminar primordium (Fig. 1a, lower panel). The embryonic labeling pattern correlates with electron microscopic observations which show that presumptive laminar cells are first contacted by incoming photoreceptor axons near the midline, and come to be displaced laterally as their cartridge matures.

The birthdates of laminar cells determined by autoradiographic analysis in the mature lamina. An analysis of the birthdates of laminar cells with respect to position in the mature lamina can be obtained by injecting \(^3\)H thymidine at a series of times during development and determining the distribution of labeled cells 4 to 5 days later, when the visual system is mature.

When embryos are injected with \(^3\)H thymidine prior to 29 hr of development, nerve cells of the mature optic lamina consistently show scattered light labeling or no label despite the presence of heavily labeled cells elsewhere in the CNS (e.g., in the supraspinal ganglion) and in nearby epithelial cells just below the carapace. More heavily labeled cells can be seen in the lamina by 29 hr, and by 30 hr the lateral and ventrolateral margins of the lamina show a bilaterally symmetric distribution of heavily labeled cells. The amount of label diminishes in cells positioned more medially and dorsally. A computer-assisted reconstruction of an adult injected at 30 hr of age is shown in Figure 3a. Heavily labeled cells (defined as cells showing 60% or more of maximum cellular label) almost exclusively occupy the lateral to ventrolateral margins of the lamina. In contrast, unlabeled neurons can be found at this position when label is injected at 32 or 33 hr of development. This can be seen in the reconstruction of an animal injected at 33 hr shown in Figure 3b. Neurons at the lateral margins of the lamina are devoid of label, whereas most cells in more medial positions are intensely labeled. Since cells at these positions fail to incorporate \(^3\)H thymidine for the duration of middle embryogenesis, we can conclude that they have withdrawn from the cell cycle by 33 hr and that an upper limit on the duration of final S phase in these cells is no more than 3 hr. Figure 3, c and d, shows similar computer reconstructions for specimens injected at 35 and 39 hr of development. Again, it can be seen that unlabeled cells occupy positions that showed label previously, demonstrating withdrawal from the cell cycle, whereas labeled cells become progressively confined to the neighborhood of the midplane. These changes can also be seen by comparing autoradiographs from adults injected with label at 35 and 39 hr of development (Fig. 2, d and e). A clear lateral-to-medial sequence of laminar cell birthdates is apparent. When \(^3\)H thymidine is given at 41 hr of age or at 43 hr (the oldest stage injected), all laminar cells are unlabeled. Many cells in the medulla of specimens injected at 41 hr are heavily labeled, as are cells in the supraspinal ganglion. Injection at 43 hr results in only a few cells labeled in the posterior medulla, whereas labeled cells remain more frequent in the supraspinal ganglion. Since laminar glial cells were not labeled in any of our series and because these cells are present when optic axons reach the lamina (LoPresti, 1975), we believe them to be the earliest cells in the lamina to become postmitotic.

Histograms of normalized grain counts over the nuclei of the lamina (Fig. 4) demonstrate the roughly 10-hr time course of cell cycle withdrawal in the lamina. Cells in the lateral lamina are not labeled by 32 to 33 hr and remain refractory to labeling for the next 10 hr. The number of unlabeled laminar cells continues to increase, and by 41
hr all laminar cells are postmitotic. The computer reconstructions indicate that birthdates of cells and their positional fates are correlated as lateral:oldest, medial:youngest. This pattern is quite similar to that of the order of arrival of ommatidial axon bundles from the eye and that of the order in which laminar cartridges are formed.

The birthdates of laminar cells within individual optic cartridges. Using serial light micrographs, it is possible to assign laminar cells to their respective cartridges with little error. From such reconstructions, the number of labeled, lightly labeled, or unlabeled cells in each cartridge can be evaluated.

A comparison of animals injected at various times during development makes it possible, in a sense, to watch the cells of each cartridge move through their final cell cycle. It is also possible to arrange cartridges in a timetable of birthdates, oldest to youngest, as presented in Table I. Several conclusions can be drawn from these analyses. First, there is an overall sequence of withdrawal of laminar cells from the cell cycle which correlates with the positions of the cells in the mature array. Second, cells of any given cartridge do not usually withdraw across all of the cartridges shows that the age of a cartridge assessed by thymidine birthdating bears a very close correlation to the time that the cartridge is formed, as observed in the embryo by electron microscopic analysis of sequential developmental stages. The arrival of photoreceptor axon bundles to the lamina and the formation of the array of optic cartridges can be viewed as a series of four steps, where a group of cartridges are formed in each step. Two cartridges are formed in the first step, and three cartridges are formed in each of three successive steps, giving a total of 11 cartridges/hemilamina (LoPresti et al., 1973). The results presented in Table I show that cartridges A, B, and C (see Fig. 1b, lower panel for the positions of these cartridges in the adult) are clearly the youngest cartridges in that some of their cells have not completed their final division several hours after the cells of the next youngest cartridge, D, are all postmitotic. This suggests that these three cartridges in the adult correspond to the last three embryonic cartridges to form at the midline. The cells of cartridges J and K withdraw from the cell cycle well ahead of the cells in the other cartridges; therefore, these two cartridges in the adult should correspond to the first two cartridges to form in the embryo. The two middle groups of embryonic cartridges should then be made up of I, G, and H as the older group and F, E, and D as the younger, more medial group. The boundary between these two groups as judged by birthdate criteria is not as sharp as the other two boundaries. These results also suggest that in the three youngest groups of cartridges (I,G,H; F,E,D; C,B,A), the most ventral cartridge should differentiate slightly before the most dorsal cartridge.

Serial electron microscopy of embryos ranging from 33 to 41 hr of development, prepared as part of the present experiments, confirm our previous observation that photoreceptor axon bundle arrivals to the lamina can be decomposed into four steps. These series also demonstrate additional intermediate stages. Figure 5 shows a summary of reconstructions from four series in which we were successful in capturing instances where only one or two of the bundles of the third group have reached the laminar primordium. The axon bundles arrive in the sequence ventral first, dorsal last. The morphological demonstration of the ventral-to-dorsal sequence of fiber bundle arrival and the accompanying degree of developmental advance within each cartridge confirm the prediction made on the basis of the birthdate results: that in the third group of cartridges, the ventral cartridge tends to be older than the dorsal one. We suspect that
this rule holds true not only for the third embryonic group of cartridges but for the following group as well, although this has not been demonstrated explicitly.

Correlation between laminar cell birthdates and photoreceptor axon bundle arrivals at the lamina. In order to establish the temporal relationship between laminar cell birthdates and the arrival of photoreceptor axon bundles at the laminar primordium and subsequent formation of cartridges, embryos were fixed at a number of times in the interval during which laminar cells are born. In some instances, it was possible to do this with the same broods used for the birthdate studies, so that embryos of exactly the same age were either injected with \[^{3}H\]thymidine or fixed for electron microscopy at injection time. The embryonic specimens were serially thin-sectioned and the number of fiber bundles reaching the lamina was counted.

The processes of fiber bundle arrival and laminar cell withdrawal from the cell cycle are contemporaneous (Fig. 6). When average rates of fiber bundle arrival and lami-
Figure 4. Normalized grain count histograms of laminar cells. Grain count data for specimens injected with \[^3^H\]thymidine at 30, 33, 35, and 39 hr of development (but fixed as adults) are displayed as normalized histograms. (Reconstructions of the laminae of these specimens are shown in Fig. 3.) Maximum cellular label was defined as the mean of the three most heavily labeled cells in each specimen and are 64, 72, 126, and 54 grains/cell, respectively, for the specimens labeled at 30, 33, 35, and 39 hr. These means were used as the normalizing factors. The histogram intervals are 10% in width.

TABLE I

The number of radioactively labeled laminar neurons in each optic cartridge for animals injected at different embryonic ages

| Cartridge | 33 | 34a | 34b | 35 | 37a | 37b | 37c | 39 | 41 |
|-----------|----|-----|-----|----|-----|-----|-----|----|----|
|           | L  | R   | L   | R  | L   | R   | L   | R  | L  |
| A         | 5  | 5   | 4   | 3  | 2   | 3   | 5   | 5  | 4  |
| B         | 5  | 5   | 5   | 4  | 5   | 4   | 5   | 5  | 4  |
| C         | 5  | 5   | 2   | 3  | 3   | 3   | 3   | 4  | 3  |
| D         | 5  | 5   | 5   | 4  | 4   | 5   | 5   | 5  | 5  |
| E         | 5  | 5   | 5   | 4  | 5   | 5   | 5   | 5  | 5  |
| F         | 5  | 5   | 3   | 5  | 3   | 4   | 3   | 3  | 3  |
| G         | 2  | 3   | 2   | 1  | 1   | 2   | 0   | 2  | 0  |
| H         | 5  | 4   | 1   | 1  | 3   | 4   | 2   | 2  | 0  |
| I         | 4  | 4   | 1   | 1  | 0   | 1   | 0   | 0  | 0  |
| J         | 0  | 0   | 0   | 0  | 0   | 0   | 0   | 0  | 0  |
| K         | 1  | 0   | 0   | 0  | 0   | 0   | 0   | 0  | 0  |
| Total Labeled Cells (%) | 79 | 74 | 49 | 50 | 51 | 56 | 53 | 52 | 17 |

*Data are given for left (L) and right (R) cartridges.
Figure 5. Intermediate stages in the arrival of the third group of photoreceptor axon bundles to the embryonic lamina. In this sequence of schematic reconstructions each number represents both the relative position of an axon bundle in the embryonic lamina and the arrival group to which that axon bundle belongs. The double vertical bar represents the dorsoventral midplane. The five cartridges formed in the first two groups of axon bundle arrivals move away from the midplane as the axon bundles of the third group arrive. The axon bundles of the third group are added sequentially in ventral-to-dorsal order. Note that the first axon bundle in the third group enters the embryonic lamina dorsomedial to the ventralmost axon bundle of the second group. The ventralmost cartridge in the second group of cartridges (the cartridges formed by the axon bundles of the second group) remains closer to the midline than do the other cartridges of the second group during subsequent development.

Figure 6. Photoreceptor axon bundle arrival to the lamina and the withdrawal of lamina cells from the cell cycle. Top, The number of photoreceptor axon bundles present in the lamina is plotted as a function of developmental age. Each point was obtained by a direct count of axon bundles from electron micrographs of a serially sectioned embryo. The circled data points at right indicate that the process of fiber bundle arrivals is complete by these times. Bottom, The number of postmitotic laminar neurons is plotted as a function of developmental age. At 30 hr all laminar cells are mitotic. At later times, the total number of postmitotic cells is approximated by considering cells with less than 10% of maximum cellular label (as previously defined) to be postmitotic. A very small error is incurred by including in this group cells that are lightly labeled. The solid circles indicate that all laminar neurons fail to label when $[^3]H$thymidine is injected at or after 41 hr of development.
series, 50 labeled cells come to be positioned in the three midline cartridges A, B, and C. The four labeled cells not found in these cartridges (the last group of cartridges to form) are located in the two dorsal cartridges adjacent to them, cartridges D and E. Of the last three cartridges to form (A, B, and C), cartridge A invariably contains the most labeled cells (four of five in every case), whereas the numbers of labeled cells in cartridges B and C vary considerably both between specimens and from side to side in each specimen. The total number of cells still in their final S phase, averaging 9 cells/hemilamina (maximum of 10 cells and minimum of 8 cells), is a conserved feature of these replicates, indeed more conserved than the distribution of these cells into cartridges. These results suggest that the rate and timing of presumptive lamina cell production are quite constant, whereas the recruitment of a particular cell into a particular cartridge is, relative to cell production, a more variable process.

Comparisons of the development of replicate animals prepared for serial electron microscopy show remarkable invariance. In the three specimens examined, seven fiber bundles have reached the lamina: the two fiber bundles of the first group, the three fiber bundles of the second group, and the two more ventral fiber bundles of the third group. A summary of the state of cartridge recruitment for the two most recently arrived fiber bundles (Fig. 7) shows no significant left-right variation, as measured by the number of presumptive laminar cells which the lead fiber of each photoreceptor fiber bundle has managed to contact. The most ventral fiber bundle had advanced deeper into the lamina than the dorsal bundle, and in two of the three specimens, the lead fiber growth cone has contacted the first cell to be recruited into its cartridge, although the characteristic "wrapping" interaction between lead fiber and presumptive laminar cell is not yet apparent. In the remaining instances, the lead fiber has only come into contact with processes of one or both of the glial cells present in the anterior lamina at this time.

The serial electron micrographs offer additional evidence that the time interval between the completion of the terminal mitoses of presumptive laminar cells and their recruitment into cartridges can be very brief, and also suggest that interaction between cells that are not yet postmitotic and the growth cones of lead fibers is possible. Posterior to the region of recruited cells in these 37-hr embryos is a region containing a number of cells of uniform size and appearance and a few cells that are mitotic. These cells are in an area which is appropriate for presumptive laminar cells that will shortly be recruited. By both position and general shape, the cells in the midline region appear to be identical to the labeled midline cells seen in embryonic light microscope autoradiographs at this stage in development. The growth cones of lead fibers extend into this region, and, in several instances, the terminal dilation of a lead fiber can be found in close apposition to mitotic cells. The correlation between photoreceptor fiber bundle arrival and cell withdrawal already indicated that very short intervals should occur between cell birthdates of laminar cells and their recruitment into cartridges. Repeated observation of lead fibers extending into the region of unrecruited, presumptive laminar cells where mitotic cells are seen confirms this. Also, lead fibers can form close appositions with a few more than the five cells they will normally recruit into their cartridge. The terminal, finger-like processes of lead fiber growth cones extend a few micrometers, bringing them into close apposition with several unrecruited cells at a time when five cells have already been recruited into the cartridge to which that lead fiber belongs.

A final point from morphological observations of the embryonic lamina concerns the stem cells of the lamina. We observe large cells, often mitotic, that may correspond to the neuroblasts described in several other arthropod visual systems (Panov, 1960; Nordlander and Edwards, 1969b) and elsewhere in the arthropod CNS (Nordlander and Edwards, 1969a; Bate, 1976). However, we have not observed an asymmetric mitosis in the immediate neighborhood of the laminar primordium, nor have we found clear columnar arrangements of what have been termed ganglion mother cells leading away from these putative neuroblasts. Analysis of the neuroblasts is further complicated by the fact that an embryonic boundary between the laminar and medullary primordia is vague.

**Discussion**

Autoradiographic analysis of the optic laminae of adult *Daphnia* labeled with [3H]thymidine during development
shows that laminar cells undergo their final mitotic divisions between approximately 30 and 40 hr of development. This occurs in an orderly sequence, such that cells located at the lateral margins of the bilaterally symmetric lamina become postmitotic first, and those cells adjacent to the midplane become postmitotic last. Thus a close correlation exists between the birthdates of cells and both the position of their cell bodies and their retinotopy in the adult array. Analysis of [\(^3\)H]thymidine labeling of the embryonic lamina, coupled with electron microscopic analysis, shows that cells already recruited into developing laminar cartridges are postmitotic. The distribution of labeled cells in embryos a few hours after injection of label demonstrates that mitotically active cells are positioned posteroomedially in the laminar primordium. Electron micrographs of the same region reveal that cells in comparable positions are not yet recruited into cartridges and are morphologically undifferentiated.

The initial contact between presumptive laminar neurons and the growing tips of lead photoreceptor axons occurs close to the midline. Newly contacted laminar cells are displaced laterally as nascent cartridges form, while in their place at the midline appear newly postmitotic cells. Occasionally, mitotic cells can be seen at the midline in close apposition with, or 1 to 2 \(\mu\)m distant from, the advancing growth cones of photoreceptor axons, indicating that the interval between final division of laminar cell precursors and the interaction of their progeny with photoreceptor axons can be quite brief. In general, a comparison of the schedules of laminar cell withdrawal from the cell cycle and arrival of photoreceptor axon bundles to the lamina indicates that the interval between cessation of DNA synthesis and photoreceptor axon contact is never more than a few hours long.

We can conclude that the interaction between presumptive laminar cells and growing photoreceptor axons, an interaction previously shown to be characterized by the envelopment of the axon by the laminar neuron and the formation of a transient gap junction between the two cells (LoPresti et al., 1973, 1974), begins as the laminar cells complete what will be their final division or within a short interval thereafter. The sequential recruitment of newly postmitotic cells at the midline, their lateral displacement as they differentiate into cartridges, and the subsequent availability of younger postmitotic cells at the midline region for newly arrived lead fibers to interact with accounts for the lateral-to-medial sequence of laminar cell birthdates.

Analysis of laminar cell birthdates within individual cartridges shows that the 11 cartridges of each hemilamina can be ordered chronologically by the birthdates of their laminar cell constituents. This temporal order agrees with the sequence of cartridge formation deduced from serial reconstructions of embryos at successive developmental stages. To maintain this ordering, the presumptive laminar cells at the midline would appear to move little relative to each other prior to their recruitment into cartridges. Following their formation, the cartridges themselves maintain a fixed array. Serial electron microscopic analysis of replicate embryos shows that the morphological sequence and schedule of developmental events is very consistent from animal to animal within a brood, as has been reported previously (LoPresti et al., 1973). Analysis of replicate autoradiographic series shows that the timing of laminar cell withdrawal from the cell cycle is also conserved. At 37 hr in development, the number of cells in the final S phase of their cell cycle varies little, but the distribution of these cells among cartridges is more variable. This suggests that the particular cartridge into which a cell is recruited is not rigidly predetermined. This suggestion is further supported by the results of experiments in which development of the lamina is indirectly perturbed by UV irradiation of the eye (Macagno, 1978). Thus, presumptive laminar cells are unlikely to be prespecified as members of a particular cartridge, from which follows that laminar cells of a single cartridge are not clonally related except by chance. Although nonclonal origin of cartridge elements cannot be decided from the evidence gathered here, it is relevant to note that there have been no studies examining clonal relations among cells in any other arthropod's optic lamina. However, genetic mosaics of the developing compound eye established either by x-ray induction in holometabolous insects or by graft transplants in hemimetabolous insects have established unambiguously that the photoreceptors of a single ommatidium have no obligatory clonal relationship (Ready et al., 1976; Shelton et al., 1977).

The synchronous withdrawal of laminar cells from the cell cycle and their innervation by photoreceptor axons raises two key questions. First, how are these two processes coordinated in time? Second, what is the significance of these events to subsequent events in the development of this neuronal array?

The first question prompts the consideration of two alternatives. Does proliferation in the laminar primordium occur independently of the development of the projection from the eye, implying that the developmental schedules of both structures are preset by some mechanism external to either structure, or does the developing eye influence proliferative events in the laminar primordium? The formation of presumptive laminar cells competent to interact with optic axons and to differentiate could be the result of an obligate and autonomous series of maturational divisions from a small number of stem cells. In the primordium of the optic ganglion of other arthropods, a progression of divisions from stem cells, called neuroblasts, to presumptive neurons has been described (Parnov, 1960; Nordlander and Edwards, 1969a). Neuroblasts can divide asymmetrically to yield neuroblasts and smaller ganglion mother cells, while the latter symmetrically divide one or more times to yield presumptive neurons. Although we have not observed such a sequence in Daphnia, the evidence does not preclude it. If it exists, such a sequence would imply an obligate series of maturational divisions leading to the production of competent presumptive laminar cells. Evidence to suggest that cell divisions generating laminar ganglion cells in the proliferative zone (the outer optic anlage in insects) are independent of the eye derives from experiments in which developing eyes of insects have been wholly or partly extirpated or in which the optic lobes have been transplanted. Although the results of earlier work are somewhat equivocal (reviewed in
Gregaria

of presumptive laminar cell production must be such

triggers, laminar cell differentiation. In general, the tim-

trophic dependence of the lamina on the eye holds true

resolution and in some cases incomplete, is also consis-

these other arthropods, much of it at a lower level of

optic laminar cartridges. The available evidence from

while the youngest ommatidia project to the youngest

ommatidia project to the oldest optic laminar cartridges,

development of the retinotectal projection in arthro-

wards, 1969b), suggests that a generalized view of the

possible exception of the butterfly (Nordlander and Ed-

and the cockroach (Stark and Mote, 1981), but with the

species within this phylum. Comparison of studies in a

which will innervate them is a feature common to all

interaction of newborn cells with the growing optic axons

and photoreceptor axon arrivals to the laminar primor-

mental rules, except of course where qualitatively differ-

this pattern of connections does not at present require

ent sorts of connections occur.

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