**Drosophila** Spectrin: the Membrane Skeleton during Embryogenesis

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**Abstract.** The distribution of alpha-spectrin in *Drosophila* embryos was determined by immunofluorescence using affinity-purified polyclonal or monoclonal antibodies. During early development, spectrin is concentrated near the inner surface of the plasma membrane, in cytoplasmic islands around the syncytial nuclei, and, at lower concentrations, throughout the remainder of the cytoplasm of preblastoderm embryos. As embryogenesis proceeds, the distribution of spectrin shifts with the migrating nuclei toward the embryo surface so that, by nuclear cycle 9, a larger proportion of the spectrin is concentrated near the plasma membrane. During nuclear cycles 9 and 10, as the nuclei reach the cell surface, the plasma membrane-associated spectrin becomes concentrated into caps above the somatic nuclei. Concurrent with the mitotic events of the syncytial blastoderm period, the spectrin caps elongate at interphase and prophase, and divide as metaphase and anaphase progress. During cellularization, the regions of spectrin concentration appear to shift: spectrin increases near the growing furrow canal and concomitantly decreases at the embryo surface. In the final phase of furrow growth, the shift in spectrin concentration is reversed: spectrin decreases near the furrow canal and concomitantly increases at the embryo surface. In gastrulae, spectrin accumulates near the embryo surface, especially at the forming amnioproctodeal invagination and cephalic furrow. During the germ band elongation stage, the total amount of spectrin in the embryo increases significantly and becomes uniformly distributed at the plasma membrane of almost all cell types. The highest levels of spectrin are in the respiratory tract cells; the lowest levels are in parts of the forming gut. The spatial and temporal changes in spectrin localization suggest that this protein plays a role in stabilizing rather than initiating changes in structural organization in the embryo.

ALTHOUGH a number of genes and gene products that specify and regulate *Drosophila* embryogenesis have been identified, understanding of developmental processes also requires knowledge of components that are mechanistic effectors of shape transformation, membrane domain formation, and cell division. Because these events in *Drosophila* are closely coordinated with the changing patterns of actin, myosin, and tubulin localization (Warn et al., 1984; Warn and Magrath, 1983; Karr and Alberts, 1986; Young, P. E., T. C. Pesacreta, and D. P. Kiehart, manuscript in preparation) and are specifically inhibited by microtubule and microfilament perturbants (Foe and Alberts, 1983; Zalokar and Erk, 1976; Warn et al., 1987), it is clear that cytoskeletal components play a major role in fly embryogenesis.

Spectrin, a cytoskeletal component that may be important in the function of membrane domains (Tokuyasu et al., 1979; Lazarides et al., 1984; Nelson and Veshnock, 1987) and that contributes to determination of shape and shape transformations in mammalian erythrocytes (Elgsæter et al., 1986), is found in many eukaryotes, including *Drosophila* (Dubreuil et al., 1987). Because spectrin may interact with microfilaments (Glenney et al., 1982), microtubules (Ishikawa et al., 1983; Sobue et al., 1987), and intermediate filaments (Langley and Cohen, 1986), and can be associated into a membrane skeleton near the cytoplasmic surface of the plasma membrane (for reviews see Bennett, 1985; Marchesi, 1985), it may be mechanistically involved in coordinating aspects of cytoskeletal function during *Drosophila* embryogenesis.

Spectrin distribution in insects has not been previously investigated, and spectrin distribution during embryogenesis is known only for mice (Sobel and Alliegro, 1985) and sea urchins (Schatten et al., 1986). As part of a broader investigation of the role of spectrin in nonerythroid cells, we have produced and characterized several antibody probes that are specific for *Drosophila* alpha-spectrin (Dubreuil et al., 1987; Byers et al., 1987). Here we use these probes to localize spectrin in developing *Drosophila* embryos and to analyze spectrin accumulation. Wherever possible we have related our observations on spectrin to other studies that have investigated the localization of actin during embryogenesis.
Our results show that Drosophila eggs contain a substantial pool of maternal spectrin that undergoes dynamic changes in distribution during early Drosophila embryogenesis. After gastrulation, new synthesis is detected and spectrin is distributed near the plasma membrane of nearly all the cells of the postblastoderm embryo. The distribution of spectrin differs, in many cases, from that of actin.

Materials and Methods

Collection of Embryos

Embryos of Drosophila melanogaster (Can-S) were grown at 25°C on grape juice–agar plates with yeast food. 1 h before placing to collection plate in a Drosophila cage, a fresh plate was introduced to stimulate deposition of any advanced embryos the females may have retained. 1 h later, this fresh plate was discarded and replaced with the actual collection plate. To harvest specific development stages the collection plate was removed from the population cage after 30-60 min and left at 25°C to develop as necessary.

Antibodies

Polyclonal rabbit serum 905 specific for Drosophila alpha-spectrin fusion protein 9a, as characterized in Byers et al. (1987), and preimmune serum collected from the same rabbit were used at 1:2,000 dilution for immunoblots or at 1:1,000 for immunocolocalization. This antisemir is referred to as polyclonal antispectrin antibody or polyclonal antibody. In most cases, the polyclonal antibody was used after affinity purification on nitrocellulose strips (Olundst, 1981) containing both paired spectrin from Drosophila S-3 cells (Dubreuil et al., 1987) or alpha-spectrin fusion protein 9a. Both preparations yield antibodies that labeled embryos in a pattern identical to the antisemir spectrum. For some experiments, monoclonal antibody M10-2, which is specific for alpha-spectrin in Drosophila S-3 cells (Dubreuil et al., 1987), was used in place of the polyclonal antibody. Culture supernatant from the clone was diluted 1:1 for use, whereas ascites fluid was diluted 1:50. Both preparations gave results which were qualitatively identical to the polyclonal antispectrin antibody in double-labeling experiments. Antibody specificity was further evaluated by preadsorbing the polyclonal antispectrin antibody to a nitrocellulose blot containing the fusion protein 9a produced from a pEV vector. In not one case did solutions that had been preadsorbed with the fusion protein produce the staining patterns described for antispectrin antibodies: only background equivalent to that seen with preimmune serum was observed. Antisemir absorbed with proteins from Escherichia coli carrying the pEV plasmid without the 9a insert gave the same staining as nonadsorbed antibody.

As an actin probe, a monoclonal antactin antibody diluted 1:100 (Lesard, 1988) was used. This antibody reacted with Drosophila actin on Western blots (data not shown). Although this antibody generally stained actin in cells, it failed, for some unexplained reason, to react with actin in the furrow canal region during cellularization. To analyze actin in this region, rhodamine phallolidin (0.33 μg/ml; Molecular Probes, Inc., Eugene, OR) was used.

Fixation for Microscopy

Embryos were placed in rinse solution (86 mM NaCl, 0.05% Tween-20, 10 mM NaNO3), dechorionated in 2.63 sodium hypochlorite (50% Clorox) (5 rain each), and either used immediately or stored at 4°C overnight.

To determine whether embryos lost spectrin during fixation, we measured the amount of spectrin lost to all preparative solutions using quantitative immunoblot. Duplicate groups of 100 embryos were collected at random from a single 10-h collection plate which included a wide range of developmental stages. The embryos were fixed as usual and then processed through all the usual solutions as for immunofluorescence (but without the addition of antibodies). All of the solutions through which the embryos had passed were combined and lyophilized. The amount of spectrin in the combined lyophilized solution residues, the fixed embryos, and an equal number of unfixed embryos was measured in qualitative immunobLOTS (see immunoblot method below). No spectrin was detected in the combined, concentrated, solution residue from 100 embryos even though the total spectrin in just one unfixed embryo contained more than the minimum amount of spectrin needed for detection. The fixed embryos retained ~70% of their spectrin immunoreactivity after having passed through all the usual preparative and staining solutions used to prepare the embryos for immunofluorescence microscopy. Thus little, if any, spectrin was lost to the solutions used during fixation and preparation for immunomicroscopy, although the spectrin retained by the embryos may have lost up to 30% of its immunodetectability as a consequence of the combined fixation steps.

Immunofluorescence

Whole mounts for immunocytochemistry were incubated in 25% goat serum in PBS for 30-60 min. Those embryos to be incubated with rhodamine-phalloidin were not exposed to the second fixative. Instead, the vitelline envelope was removed with tungsten needles, and the embryos were put back into the aqueous component of the first fixative for 1 h.

For cryosectioning, embryos were used following the two-stage procedure, incubated in gradually increasing concentrations (10, 30, 50%) of enzme grade sucrose (Bethesda Research Laboratories, Gaithersburg, MD) in PBS at 4°C over several hours, mounted in O. C. T. (Miles Laboratories, Naperville, IL), frozen in dry ice, sectioned at a thickness of 5 μm on a Microsome (International Equipment Co., Needham Heights, MA), and mounted on glass slides coated with 1% gelatin dissolved in 0.1% chromium potassium sulfate.

Nomenclature

We adhered to the terminology of Foe and Alberts (1983) to describe the nuclear cycles: each nuclear cycle begins with the start of interphase and ends with the conclusion of mitosis. The timing given for each stage is that observed under our conditions.

When describing structures near the plasma membrane, we used the perspective of the plasma membrane being "above" the cytoplasm. "Apical" refers to the region of each cell that is or was nearest to the embryo surface during the cell cycle. The study has analyzed the distribution of only the alpha chain of spectrin. Thus, "spectrin" refers only to alpha-spectrin. Work using antibodies against the beta chain of Drosophila...
spectrin has been initiated, but there is no evidence that the two chains are
differently localized in the early embryo.) "Caps" refer to concentrations of
spectrin or actin that are associated with the plasma membrane.

**Quantitative Immunoblots**

The total amount of spectrin per embryo was measured at different stages
during embryogenesis. Because no system of collection was available that
adequately synchronized development among populations of embryos, em-
bryos were individually examined with a dissecting microscope to deter-
mine their stage of development. The following stages were defined by con-
venient morphological criteria that could be visualized in a dissecting
microscope. The approximate age of the embryo (indicated in parentheses)
is an estimate, based on normal development at 25°C (Campos-Ortega and
Hartenstein, 1985): (a) pole cells absent (0–60 min); (b) pole cells present
(60–120 min); (c) ongoing cellularization or amnioproctodeal invagination
in external position (120–220 min); (d) amnioproctodeal invagination not
visible, with the cephalic furrow the major external feature (220–260 min);
(e) anterior segmentation evident (320–440 min); (f) segmentation promi-
ient over almost the entire length of the embryo, with germ band shortening
(440–560 min); (g) clypeolabral shortening (560–620 min); and (h) midgut
folded into a linear array of segments (780–960 min).

Embryos were dechorionated in 50% bleach and placed in rinse solution
for microscopic inspection. Three samples, staged as described above, with
eight embryos per sample, were collected for each developmental stage.
Homogenization buffer was prepared by placing 20 μl of 5 × SDS-PAGE

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**Figure 1.** Spectrin in preblastoderm embryos. (a) Whole mount of a nuclear cycle 2 embryo, parasagittal optical section. Spectrin fluores-
cence is visible at the plasma membrane (arrow) and throughout the central regions of the embryo. (b) Whole mount of a nuclear cycle
7 embryo, optical section, labeled with preimmune serum (photographic conditions identical to c). (c) Spectrin fluorescence from whole
mount of nuclear cycle 7 embryo, optical section, double labeled with antispectrin serum and Hoechst 33258. Because the optical section
shown is not parasagittal, spectrin fluorescence at the plasma membrane is not visible in the photograph, although spectrin fluorescence
was evident at the plasma membrane and is seen concentrated in cytoplasmic islands (arrows) that surround nuclei (shown in d). (d) Same
as c, but illuminated to visualize Hoechst 33258 staining. The nuclei are grouped near the longitudinal axis of the embryo in a pattern
that resembles that of the spectrin-containing cytoplasmic islands shown in c (arrows). (e) Same embryo as a, at higher magnification.
Parasagittal optical section of the plasma membrane and associated spectrin (arrow). (f) Surface view of embryo in a. Bars: (a–d) 100
μm; (e and f) 10 μm. (Except where serum is specified, all of the figures in this paper show fluorescence due to binding of affinity-purified
antispectrin antibody. Binding was detected with secondary antibody as described in Materials and Methods.)
During nuclear cycle 9, spectrin becomes concentrated into caps near the plasma membrane (Fig. 2 a), comparable to the caps of actin that also overlie the nuclei (Karr and Alberts, 1986). A few embryos have spectrin caps at anaphase 9, others at telophase 9 (Fig. 2 b), and all do by the end of interphase 10. Many of the caps form as the nuclei divide eight times, beginning their movement to the surface at the end of nuclear cycle 7 (for a description of embryogenesis see Campos-Ortega and Hartenstein, 1985). During this period, spectrin is concentrated subjacent to the plasma membrane (Fig. 1, a and b), in microtubule-rich regions (Karr and Alberts, 1986) that surround nuclei in the interior of the early syncytial embryo (Fig. 1, c and d), and, more diffusely, throughout the remaining embryo cytoplasm (Fig. 1, a, c, and e). Although spectrin appears as a continuous band near the plasma membrane in sagittal optical sections (Fig. 1 e), face views show that it forms an apparently random reticulum near the entire surface of each embryo (Fig. 1 f). The distribution of spectrin shows no evidence of being correlated with the radial migration of the nuclei from the center of the embryo to the surface and is not obviously polarized with regard to the dorsi-ventral or anterior-posterior axes of the early embryo. At a variable time during nuclear cycle 9, spectrin begins to redistribute into caps near the plasma membrane (see below).

Nuclear Cycle 9 through Interphase of Nuclear Cycle 10 (from ~70 to 85 min): Spectrin Caps Form

During this period, the somatic nuclei complete their migration from the inner regions of the embryo and come to lie within ~2.5 $\mu$m of the plasma membrane. Interphase of nuclear cycle 10 marks the beginning of the syncytial blastoderm period. During nuclear cycle 9, spectrin becomes concentrated into caps near the plasma membrane (Fig. 2 a), comparable to the caps of actin that also overlie the nuclei (Karr and Alberts, 1986). A few embryos have spectrin caps at anaphase 9, others at telophase 9 (Fig. 2 b), and all do by the end of interphase 10. Many of the caps form as the nuclei are in the process of approaching the embryo surface (Fig. 2 b). Although the timing of cap formation varies from em-
bryo to embryo, the movement of spectrin into caps must be abrupt: partially formed spectrin caps are rarely seen, and <15 min elapses between the beginning of anaphase 9 and the end of interphase 10 (Foe and Alberts, 1983). In face view, each cap is circular or sometimes slightly elliptical (Fig. 2 a), with an average diameter of 15 \mu m \( (n = 10; \text{ SD } = 1.7 \mu m) \). A nucleus or mitotic figure is located in the cytoplasm below each cap (Fig. 2 b). In sagittal optical sections (Fig. 2 c), the amount of spectrin between the caps appears to be less than was present before cap formation, as though preexisting spectrin had relocated and concentrated to form the caps (compare Fig. 1 e with Fig. 2 c).

Antispectrin and antiactin antibodies were used in double-labeled embryos to evaluate the association of spectrin and actin. Before nuclear cycle 9, actin is present throughout the embryo at low levels and is concentrated uniformly near the plasma membrane, as previously described (Karr and Alberts, 1986; Warn and Magrath, 1983). Our results show that actin caps form before the time reported by Karr and Alberts (1986), appearing between anaphase 9 and early interphase 10. Although spectrin caps begin to form at this time, small spectrin caps are consistently present before spectrin cap formation (Fig. 3, a and b). Once formed, the two proteins codistribute within each cap (Fig. 3, c and d).

**Nuclear Cycles 10–13 (from ~85 to 130 min): Spectrin Caps Elongate, Separate, and Collide with Other Caps**

Although pseudo cleavage furrows form at each prometaphase during nuclear cycles 10–13, these furrows are transient and do not lead to cellularization (Stafstrom and Staehelein, 1984). During prophase 10, the centrosomes divide and move toward the eventual position of the spindle poles. Anticipating the division and directional movement of the chromosomes, the spectrin caps elongate during interphase and prophase, and the central portions of the elongated caps become narrowed (Fig. 4, a and b), presumably in the regions where daughter caps will separate from each other during the subsequent metaphase and anaphase (Fig. 4, c and d). Spectrin concentration in the region of incipient cleavage is no greater than in other regions of the cap. Thus, cap division does not appear to involve a constrictive or compressive mechanism, but rather an oppositely directed movement of daughter caps. During anaphase and telophase, the distance between daughter caps increases (Fig. 4 e) and their lineage becomes obscure. Inspection of the underlying Hoechst-stained telophase figures (Fig. 4 f) shows that in many cases the daughter caps collide with adjacent, nondaughter caps forming what appear to be slightly compressed areas of high spectrin concentration (Fig. 4 e). By interphase 11, the newly
formed nuclei are generally centered under each cap. The apparent linkage between spectrin cap elongation, cap division, and karyokinesis is maintained during the next three rounds of mitosis, but becomes increasingly difficult to follow as the number of nuclei and caps increases and the distinction between cap and intercap regions is blurred. Although Warn et al. (1984) and Karr and Alberts (1986) report that the actin caps spread out and move into the pseudo furrows that form during the prophase-metaphase period of nuclear cycles 11-13, at no stage did we detect increased concentrations of spectrin near the pseudo cleavage furrows.

**Interphase, Nuclear Cycles II-13, and Pole Cell Development: Spectrin Is Concentrated in the Plasma Membrane and the Subjacent Cytoplasm**

As the number of nuclei in a syncytial blastoderm stage embryo increases, nuclear size decreases (Zalokar and Erk, 1976). During interphase, the plasma membrane above each nucleus curves to produce an array of domes (Turner and Mahowald, 1976). Optical sections obtained with the scanning confocal microscope show that the distribution of spectrin follows the contour of the plasma membrane. Spectrin caps are no longer distinguishable, and spectrin fluorescence
Figure 5. Syncytial blastoderm during interphase. (a–d) Successive scanning confocal microscope optical sections through part of a nuclear cycle 12 whole mount embryo. The optical sections are ~2 μm thick, and are taken at 4-μm increments, starting from the embryo surface in a. Arrows in b point to membrane-associated spectrin. Spectrin fluorescence is also clearly visible in the cytoplasm surrounding the nuclei (N). The concentration of spectrin in the cytoplasm diminished farther into the embryo (c and d) where the outlines of the nuclei and some yolk granules are difficult to detect. (e) Scanning confocal microscope optical section within a nuclear cycle 9 embryo whole mount. Spectrin is concentrated at the plasma membrane of pole buds (PB). (f) Cryosection of late syncytial blastoderm embryo. At this late syncytial stage, spectrin is present at the plasma membrane above the somatic nuclei (N), but is barely detectable at the plasma membrane of pole cells (PC). Bars: (a–d) 10 μm; (e) 7.5 μm; (f) 10 μm.

is uniform in en face optical sections of the surface of the embryo (Fig. 5 a). 4 μm farther into the same embryo, fluorescence encircles the nuclei of the syncytial blastoderm, but is greatest where the plasma membrane extends into the cytoplasm at the perimeter of each dome of plasma membrane (Fig. 5 b). Lesser concentrations of spectrin are evident in the cytoplasm between the unstained nuclei and the intensely stained plasma membrane (Fig. 5, b and c). At this stage of development, little or no spectrin is detectable near the yolk granules toward the center of the embryo (Fig. 5 d).

During nuclear cycle 9, spectrin distribution in the pole buds is similar to that found in the somatic regions (Fig. 5...
Figure 6. Cellularization. (a) Sagittal optical section through an embryo whole mount during the first (slow) stage of cellularization. Spectrin is concentrated near the furrow canals (arrowhead) which at this stage are close to the embryo surface (arrow). (b) Same as in a, but during the second (fast) stage of cellularization. The furrow canals have just passed the base of the elongated nuclei. The concentration of spectrin is highest just behind the advancing furrow canals (see Fig. 7, colocalization study with actin) with little spectrin at the embryo surface. (c) Cryosection of an embryo at a stage of cellularization similar to, or slightly later than, that in b. Spectrin is concentrated in the region near the furrow canals. (d) Cryosection of embryo in which cellularization is almost completed. The relative concentration of spectrin is now highest at the embryo surface, whereas spectrin near the furrow is almost undetectable. Bars: (a and b) 20 µm; (c and d) 10 µm.

Later, during the syncytial blastoderm stage, the amount of spectrin near the plasma membrane of pole cells begins to decrease (Fig. 5 f). Our observations of living embryos (Kiehart, D. P., P. E. Young, and S. Inoue, unpublished observations) indicate that the timing of this spectrin decrease varies somewhat from embryo to embryo, but, in general, coincides with a period of increased movement and microspike formation at the pole cell surfaces (not shown).

**Cellularization (from ~130 to 170 min): Spectrin Shifts between the Furrow Canal and the Embryo Surface**

After nuclear mitotic cycle 13, a distinctive period of cellularization occurs: each of the 5,000 or so cortical nuclei of the syncytial blastoderm, together with surrounding cytoplasm, is enclosed by plasma membrane to form the individual epithelial cells of the cellular blastoderm. This period is accompanied by striking changes in spectrin distribution. Immediately before cellularization, spectrin is uniformly concentrated near the plasma membrane. But during the first, slow phase of furrow growth (Fullilove and Jacobson, 1971), when many new microspikes appear at the embryo surface (Turner and Mahowald, 1976), the distribution of spectrin changes abruptly and becomes most concentrated near the furrow canal (Fig. 6 a). F-actin is also associated with the furrow canals, although a meshwork of actin filaments remains close to the cell apices (Warn and Magrath, 1983).

High concentrations of spectrin remain associated with the furrow until at least the beginning of the second phase of furrow growth, as the furrow proceeds deeper into the embryo and passes the elongated nuclei (Fig. 6, b and c). By this stage, there is less actin at the cell apices, and the F-actin associated with the developing lateral cell membranes is more densely packed (Warn and Magrath, 1983). Double-labeled preparations show that the region of highest actin concentration at the furrow canal is always slightly in advance of the most concentrated region of spectrin (Fig. 7, a and b). Late during this second phase of furrow growth, spectrin fluorescence almost disappears from the region near the furrow canal and becomes increasingly evident near the embryo surface (Fig. 6 d). Thus, within the 40-min period of cellularization, spectrin appears to cycle from the embryo surface, to the growing furrow, and then back again to the embryo surface. The return of spectrin to the embryo surface appears to coincide with the time during which the embryo surface becomes smooth (Turner and Mahowald, 1976) and the actin networks at the furrow tips become distinct, presumably contractile, rings (Warn and Magrath, 1983).

**Gastrulation (from ~170 to 190 min): Spectrin Is Concentrated in the Apical Regions of the Epithelial Cells**

Gastrulation is initiated during the terminal phase of cellularization. The relative concentration of spectrin near the embryo surface continues to increase during the early stage of gastrulation (Fig. 8 a), and immunofluorescence is brightest in distinct regions of the embryo, such as the developing amnioproctodeal invagination and the cephalic furrow (Fig. 8 b). To determine if these localized bright spots result from cell superposition, we immunostained 5-µm-thick cryosections of embryos that contained only a monolayer of cells. In these sections it is apparent that the cells of both the amnioproctodeal invagination and the cephalic furrow do indeed contain relatively high concentrations of spectrin in their apical regions (Fig. 8 c). In contrast, the pole cells showed lower levels of spectrin (Fig. 5 f and Fig. 9) where
double-label experiments (Fig. 9, a and b) clearly showed the presence of actin.

The localization of spectrin at the apical membranes of somatic cells during late cellularization and early gastrulation is in striking contrast to the more generalized distribution of spectrin around almost every cell in the embryo after just 1 h of further development (see below).

**Postblastoderm Mitoses, Segmentation, and Later Development (from ~190 to 960 min): Spectrin Distribution Becomes Generalized**

The end of gastrulation marks the beginning of the germband elongation stage. During the ensuing postblastoderm period of cell proliferation and tissue formation, spectrin becomes evenly distributed around the periphery of most cells (Fig. 10 a), but in amounts that vary from tissue to tissue. Thus, the spiracles and tracheal pits of the developing respiratory system contain high concentrations of spectrin (Fig. 10 b), whereas spectrin is difficult to detect in cells of the proventricular structure which joins the segments of the developing gut (Fig. 10 a). In addition to the peripherally localized spectrin, low concentrations of spectrin are diffusely distributed throughout the cytoplasm of both interphase and mitotic phase cells (Fig. 10, c and d). Although yolk granules are a major portion of the embryo, they never contain spectrin (Fig. 10 a).

**Spectrin Accumulation during Embryogenesis**

When immunostained embryos of different developmental stages are photographed together, it is evident that early embryos, including gastrulae, contain lower levels of spectrin immunofluorescence than do segmented embryos after germband shortening. Confocal optical sections through side-by-side embryos (Fig. 11 a) show that the greater fluorescence in the more mature embryos cannot be attributed simply to the superposition of cells. Rather, these optical sections suggest that the older embryos contain a greater amount of spectrin per unit surface area of plasma membrane than do the younger embryos.

Developmental immunoblots of whole embryo homogenates were used to quantify the changes in the level of spectrin during early embryogenesis. During the initial stages of development, up to and including gastrulation, the amount of spectrin per embryo remains approximately constant (Fig. 11 b). Immediately after gastrulation, as the cephalic furrow becomes the major visible morphological feature, the amount of spectrin begins to increase. The total amount of spectrin reaches a plateau as germband shortening occurs and segmentation becomes evident along the length of the embryo. This overall pattern of accumulation was duplicated in each of several experimental collections, but the proportional increase in spectrin varied between collections, with the amount of spectrin in late stage embryos being between 3.5- and 5.5-fold that found in the gastrula and pregastrula stages.

Using a known concentration of Drosophila alpha-spectrin as a standard, we calculate that late stage embryos contain on average 4.5 ng of alpha-spectrin per embryo. Assuming that such an embryo contains 40,000 cells and that the spectrin is equally distributed among them, each cell would contain \( \sim 10^{-4} \) ng of alpha-spectrin. This represents \( \sim 3 \times 10^8 \) chains of alpha-spectrin/cell, a number remarkably close to the number of alpha chains in the \( 2.4 \times 10^9 \) molecules of spectrin heterodimer in mature mammalian erythrocytes (Agre et al., 1985). A mammalian erythrocyte is comparable in size and surface area to a cell of the late stage Drosophila embryo.

**Discussion**

Our results show that during early Drosophila embryogenesis major events such as cellularization and gastrulation are
accompanied by striking, systematic changes in spectrin localization. Because the major formative events of early development occur during a period when the total amount of spectrin in an embryo does not increase, we infer that the accompanying localized changes in spectrin concentration in areas such as caps, furrow canals, or the apical regions of the gastrulating epithelial cells are probably the result of redistribution rather than de novo synthesis. This conclusion is consistent with the observation that the localized changes in spectrin concentration are rapid and that the accumulation of spectrin at one location often coincides with the diminution of spectrin at another. For example, the concentration of spectrin into caps coincides with a reduction of spectrin between the caps (compare Fig. 1 e with Fig. 2 c), and the deposition of spectrin near the apical surface of newly formed epithelial cells coincides with the diminution of spectrin at the basolateral regions of these cells (Fig. 6 d).

Observations of total spectrin during embryo development (Fig. 11) suggest that some of the developmental changes in spectrin localization involve the recruitment of unassembled maternal spectrin to the assembled state. Given the size of a Drosophila embryo and the average size of its cells, simple calculations show that the three rounds of postblastoderm mitoses that result in some 40,000 new cells must increase the total plasma membrane surface area at least 20-fold in comparison with the plasma membrane surface area in the syncytial embryo. Because the increase in membrane surface area substantially exceeds the 3.5–5.5-fold increment in total embryo spectrin (Fig. 11 b), the enhanced spectrin immunofluorescence at the plasma membrane of nearly all the cells produced by these postblastoderm mitoses (Fig. 11 a) cannot be accounted for exclusively by the increase in total spectrin. We therefore hypothesize that significant amounts of diffusely distributed, preexisting maternal cytoplasmic spectrin are recruited to the membrane during postblastoderm mitoses. If this hypothesis is correct, it implies that maternal spectrin can continue to play an important role in Drosophila development past the stages of embryogenesis examined here. We have begun to characterize several putative spectrin-deficient mutants. Their survival to the stage of third instar larvae is consistent with the sustained influence of maternal spectrin.

If diffusely distributed cytoplasmic maternal spectrin is recruited to cell surfaces and accounts for much of the fluorescence associated with the 20-fold increase of the plasma membrane surface area of postblastoderm embryonic cells, then cytoplasmic spectrin in the early syncytial stages (Figs. 1, 4, and 11) must represent a large fraction of embryonic spectrin. While spectrin is frequently defined as a plasma membrane-associated protein, our observations in Drosophila, as well as other observations in mammalian cells (Mangeat and Burridge, 1984; Nelson and Veshnock, 1987; Black et al., 1988), indicate that there can be important amounts of spectrin that are not associated with the cell surface. The diffuse appearance of this cytoplasmic immunofluorescence in comparison with the localized immunofluorescence at the embryo surface may easily lead to underestimates of the proportion of total spectrin in the cytoplasm.

Although spectrin caps appeared after actin cap formation, the spectrin caps always colocalized with the actin caps above the early syncytial–stage nuclei. Thus, spectrin was seen to concentrate in cytoplasmic domains that could be...
Figure 9. The forming amnioproctodeal invagination. Optical section in whole mount embryo double labeled for spectrin and actin. Pole cells (arrows) and the embryo surface (arrowheads). (a) Spectrin (antispectrin serum) is concentrated at the surface of the invagination and along very short portions of the lateral plasma membrane of each cell. Little spectrin is in the pole cells. (b) Actin (anti-actin antibody) is diffusely present near the surface of the embryo, including the amnioproctodeal invagination where spectrin is concentrated (see Fig. 8 c). Actin is more concentrated in the pole cells. (c) Nomarski micrograph of area in a and b. Bar, 10 μm.

Figure 10. Late stages of development. (a) Cryosection of embryo interior showing spectrin (antispectrin serum) associated with many cell types except those of the proventriculus (arrow); control sections without antibody show that the area in the center of the proventriculus is autofluorescent (not shown). No spectrin is present in the yolk granules (Y). (b) Whole mount showing spectrin concentrated in the region of the tracheae (arrows). (c) Whole mount, optical section near the embryo surface, showing prophase cells (arrows) in which spectrin is diffusely distributed throughout the cytoplasm; arrowheads indicate interphase cells in which spectrin is clearly absent from the nuclei. (d) Hoechst staining of embryo in c showing prophase figures (arrows) and interphase nuclei (arrowheads). Bars: (a) 10 μm; (b) 100 μm; (c and d) 10 μm.
presence of spectrin stabilizes structure, its removal should therefore reason that spectrin may frequently sustain or preceded other cytoskeletal changes during embryogenesis. Actin (and myosin) concentration. These and other examples showed that spectrin localization often succeeded rather than preceded other cytoskeletal changes during embryogenesis. We therefore reason that spectrin may frequently sustain or stabilize organization rather than initiate reorganization in Drosophila embryogenesis. We reason further that if the presence of spectrin stabilizes structure, its removal should facilitate structural change. Thus, the decrease of spectrin at the plasma membrane of pole cells during the syncytial blastoderm stage and the withdrawal of spectrin from the embryonic plasma membrane toward the lateral faces of the furrow canal during early cellularization may contribute to the simultaneous enhancement of membrane activity, such as microspike formation, that coincides with the diminution of spectrin at these membrane surfaces.

Despite spectrin’s widespread occurrence and remarkably conserved structural features (Marchesi, 1985; Byers et al., 1987), its role in nonerythroid cells remains a puzzle. Although there is evidence that spectrin may maintain the physical state of the plasma membrane, modulate membrane lipid organization, stabilize membrane domain composition, or link cytoskeletal elements to the plasma membrane (Baines, 1984; Bourguignon et al., 1985; Harris et al., 1986; Nelson and Veshnock, 1986; Del Buono et al., 1988; Lee, J. K., J. D. Black, E. A. Repasy, R. T. Kubo, and R. B. Bankert, manuscript submitted for publication), one of the few experiments that attempted to directly assess spectrin’s cellular function in nonerythroid cells did not produce data that supported an obvious role for this protein (Mangeat and Burridge, 1984). Our immunofluorescence studies in Drosophila extend knowledge of spectrin’s localization to an invertebrate system in which a combination of genetic, molecular, and cell biological approaches should make it possible to clarify the precise functions of this protein.

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