The Structure of Cytoplasm in Directly Frozen Cultured Cells. I. Filamentous Meshworks and the Cytoplasmic Ground Substance

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ABSTRACT Cultured fibroblasts or epithelial cells derived from *Xenopus laevis* embryos were directly frozen, freeze-substituted by an improved method, and then either critical-point-dried and viewed as whole mounts, or embedded and thin sectioned. In thin regions of these cells, where ice crystal artifacts are absent, the cytoplasm consisted of a dense, highly interconnected meshwork of filaments, embedded in a finely granular ground substance. The meshwork in directly frozen, intact cells was compared with that in cells that were lysed (physically, with detergents, or with filipin), or fixed with glutaraldehyde before freezing. Although filaments tended to be less numerous in lysed cells, their overall organization was the same as that in intact cells. However, fixation with glutaraldehyde before freezing distorted the meshwork to variable degrees depending on the osmolarity of the fixation buffer, and also obscured the granular ground substance which is obvious in directly frozen cells. With optimal preparative methods, the cytoplasm of these directly frozen cells is shown to consist of a cytoskeleton composed of discrete interwoven filaments interconnected by numerous finer filaments and a readily extractable granular matrix which presumably represents aggregations of cytoplasmic proteins.

Electron microscopic examination of detergent-extracted, fixed cultured cells prepared by negative staining (34), as whole mounts (29, 30), or frozen, etched, and prepared as platinum replicas (9) has contributed greatly to an understanding of filamentous meshworks in their cytoplasm. However, the relationship of these meshworks to soluble cytoplasmic components, and even the extent to which preparative procedures affect the organization of filaments, remains unclear.

It is readily apparent from the recent literature that different preparative procedures and methods of viewing cells in the electron microscope have led to various interpretations of cytoplasmic fine structure. These interpretations range from those based on high-voltage electron microscopy of whole mounts of cultured cells, (4, 23, 38) to those derived from replicas of rapid frozen cytoskeletons prepared by detergent extraction (9), or from detergent-extracted, negatively stained cultured cells (33). One view is that the cytoplasmic ground substance or matrix is made up of a three-dimensional network of slender strands (microtrabecula) that represent a polymerized, protein-rich phase separated by water-rich solutions of low molecular weight components (23). The microtrabecula are presumed to have a structural role, interconnecting known cytoskeletal elements such as stress fiber filaments and microtubules giving the cytoplasm the form of a structured gel. An alternative view is that the ground substance consists of a randomly distributed amorphous granular material, which presumably represents a protein sol (9, 33). Proponents of this view emphasize that the cytoskeleton appears to be primarily composed of discrete filaments of known composition (F-actin, intermediate filaments, and microtubules). Further elaborations on these different interpretations have appeared more recently, but disagreement persists (12, 16, 22).

It is clear from these previous results that the discreteness of the filaments and their relationship to the cytoplasmic ground substance is the major area of controversy. Therefore, we have attempted to examine the contents of intact cells by
combining the advantages for structural preservation offered by direct freezing and freeze-substitution with the appreciation of structure in these dimensions afforded by examining whole mounts in high-voltage electron microscopes. Although some previous data using whole mounts after freezing and freeze-substitution is available (22, 38), our efforts to improve freezing and freeze-substitution appear to provide a substantially more realistic view of cytoplasm. This improvement justifies a new and detailed look at cytoplasmic fine structure.

In this paper, we concentrate on the relationship between filamentous meshworks and the ground substance in which they are suspended. The regional organization of the cytoplasm, as determined by patterns of organelle and cytoplasmic movement, is described separately (2). Our interpretation of cytoplasmic fine structure differs in several important ways from those of previous studies using intact or extracted cells prepared by either fixation or direct freezing.

MATERIALS AND METHODS

 Cultures: Dissociated cell cultures were derived from the somites or ectoderm of Xenopus laevis. Cultures from somites contained muscle cells and fibroblasts, while cultures from ectoderm contained fibroblasts and epithelial cells. Details of the culture methods have been published elsewhere (3, 21). Dissociated cells were pipetted onto gold or titanium grids coated with Formvar or carbon while immersed in culture medium (88% Steinberg's solution, 10% L-15, 1% fetal bovine serum, 10,000 IU penicillin/streptomycin; osmolarity: 150 mOsmol). Glow-discharged or polylsine-coated carbon films on 300-mesh titanium grids were preferred because they were the most electron lucent and stable under the electron beam.

Albumin Mounts: Chicken egg albumin (Sigma Chemical Co., St. Louis, MO) solutions prepared in concentrations of 2.5% in distilled water served as controls for freezing and other preparative steps. Coated grids were touched to the surface of a drop of the solution until enough liquid to cover one third to one half of the grid spread onto the grid surface. The grid was then frozen immediately.

Detergent and Filipin Extraction: Cells subjected to detergent or filipin extraction were first rinsed for 15 s in extraction buffer (30) that contained 30 mM Pipes, 25 mM HEPES, 10 mM EGTA, 1.7 mM MgSO4 and which was neutralized to pH 6.9 with KOH. The osmolarity of the buffer was 140 mOsmol. Cells were then lysed with extraction buffer supplemented with either 0.1% Triton X-100, 0.1% saponin, or a mixture of 0.04% filipin and 1% DMSO for 30–60 s. Lysis was terminated by freezing.

Freezing: Cultures on grids were frozen by rapid injection of the grid into a rapidly stirred mixture of propane/isopentane (3:1) (13) or propane/ethane (3:1) cooled by liquid nitrogen. Injection of the grids into the freezing mixture was accomplished by means of a spring-driven device (6, 8). Rapid stirring of the freezing mixture and careful removal of excess fluid from the grid immediately before freezing was critical to obtaining good freezing. Because the culture medium was rather viscous, grids were rinsed in Steinberg's solution (osmolarity: 134 mOsmol) before freezing. Some cultures grown on grids were also frozen by slamming them against a copper block cooled by liquid helium (10).

Freeze-Substitution: While under liquid nitrogen, grids to be freeze-substituted were placed in a stainless-steel holder, which was then placed in a scintillation vial containing 10 ml of the primary substitution fluid covered with 10 ml of liquid nitrogen. The vial was usually transferred to a freezer maintained at −80°C to −78°C, though passive warmup in a well-insulated box was also successful. Temperature was monitored by a digital thermometer and scintillation vial containing 10 ml of the primary substitution fluid covered was neutralized to pH 6.9 with KOH. The osmolarity of the buffer was 140 mOsmol. Cells were then lysed with extraction buffer supplemented with either 0.1% Triton X-100, 0.1% saponin, or a mixture of 0.04% filipin and 1% DMSO for 30–60 s. Lysis was terminated by freezing.

Electron Microscopy: Most of the electron microscopy for this study was performed on a JEOL-200 CX with a high magnification pole piece operating at 200 kV. In some instances, a tilting liquid nitrogen cold stage (Gatan, Inc., Warrendale, PA) was used to evaluate heating effects caused by the electron beam. We also used the JEOL 1,000-kV high-voltage electron microscope at Boulder, Colorado for parts of this study. Filament diameters were measured in high magnification micrographs (×500,000) calibrated with a waffle grating using a digitizer connected to a desk-top Hewlett-Packard computer (Hewlett-Packard Co., Salt Lake City, UT).

RESULTS

Selection of Optimal Preparations

The cytoplasm of well-frozen cells appeared uniformly dense at low magnifications (Fig. 1) but at higher magnifications was finely textured (Fig. 2). Poorly frozen cells (not shown, but see Fig. 3) showed a distinct reticular pattern at low magnifications and at higher magnifications appeared to have a coarse texture that resulted from irregular-shaped spaces in the cytoplasm. The sizes of the irregular-shaped spaces in the cytoplasm of poorly frozen cells varied and were presumably dependent on the sizes of the ice crystals that were formed during freezing. Although poor-quality freezing was easily detectable, it was more difficult to make a precise judgment about marginal freezing quality.

To simplify interpretation of ice crystal–induced artifacts we divided the quality of freezing into four categories: (a) Poorest quality—large ice crystal–induced spaces (80 nm); membranes distorted, filaments grossly distorted or even unrecognizable. (b) Poor quality—large filaments and microtubules recognizable but distinct ice crystal–induced spaces (40–50 nm) were uniformly distributed throughout the cytoplasm. Fine filaments and ground substance were distorted and halos were found around membrane-limited organelles. (c) Fair quality—large filaments were undistorted, small filaments were undistorted in some areas. Ice crystal–induced fissures (20–30-nm width) traveled through the cytoplasm, distorting ground substance and fine filaments. Areas between fissures might appear free of ice crystals. Halos were found around some organelles. (d) Good quality—very few spaces <5 nm detected which were attributable to ice crystals. All filaments appeared distinct and of uniform diameter along their lengths. Ground substance appeared uniformly dense with an ex-
tremely fine granular texture (Figs. 1 and 2).

Well-spread fibroblasts or epithelial cells had relatively thick organelle-packed central regions surrounded by extensive peripheral regions that were usually very thin and contained few organelles (2). Precise assessments of freezing quality as well as the majority of the observations presented in this paper were made from the thin peripheral regions of cells. For assessments of freezing quality it was useful to compare whole mounts with thin sections because the latter gave the information in a form that was more easily interpreted and familiar.

Solutions of pure chicken egg albumin were also frozen and processed in the same way as the cell cultures. Frozen, freeze-substituted, and critical-point-dried albumin solutions (2–5%) showed a range of ice crystal–induced spaces similar to those in the ground substance of whole mount cells. Filament-like structures also formed in the albumin when freezing was less than optimal. Their exact appearance varied depending on the size of the ice crystal–induced spaces (Fig. 3, B and C).

Optimal frozen albumin had a homogeneous, finely granular appearance (Fig. 3A). The best quality of freezing obtained by immersion in propane mixtures was equivalent to the best quality obtained by slam-freezing against a pure copper block cooled by liquid helium.

Artifacts associated with electron beam damage could also be seen in whole mount cells, especially those that were not coated with a layer of carbon. These artifacts ranged from stretching, tearing, or sometimes folding and shrinkage of the cytoplasmic matrix or dense organelles, to a barely noticeable clearing of the cytoplasm. Viewing uncoated cells at −160°C using a liquid nitrogen cold stage (at 200 kV) slowed but did not prevent electron beam damage. Carbon coating could also help alleviate such effects but it was still necessary to minimize the electron beam dose.

Artifacts associated with critical-point drying have been described recently (25, 26). We initially observed similar artifacts when residual water or intermediate fluid (acetone) was not completely removed from the critical-point-drying
chamber. Cells properly critical-point-dried were the same as those that were freeze-dried. Dried preparations had to be stored in a very dry environment. Even very short exposures (1 min) to normal atmosphere resulted in drying artifacts.

**FIGURES 2 and 3** Fig. 2: Higher magnification stereo view from the cell shown in Fig. 1. The area shown is a thin, peripherally located area of the filamentous meshwork. Granules are packed between filaments. The insets show at higher magnification examples of the different size classes of filaments: 9–14, 7–8, and 4–6 nm. Filaments are indicated by the arrowheads. 1,000 kV. × 48,000. (Inset) × 77,000. Fig. 3: Comparison of three different regions of preparations of directly frozen chicken egg albumin that was freeze-substituted in acetone with a sequential fixative combination of acrolein-tannic acid/osmium/glutaraldehyde followed by critical-point drying. (A) Optimal freezing; (B) fair freezing; (C) poor freezing. Arrows indicate spaces induced by ice crystals. 200 kV. × 110,000.

**Directly Frozen, Freeze-substituted Cells**

Stereo views of thin peripheral regions of cells revealed a dense, interconnected, three-dimensional meshwork of fila-
ments (Fig. 2). Long, slightly curving filaments that sometimes ran in loose bundles, and interconnecting short, straight filaments that often formed Y-shaped junctions accounted for most of the filaments in the meshwork. The loose bundles of long, curving filaments were occasionally continuous with the tightly packed bundles of stress fiber filaments. Long, curving filaments were usually 7-8 nm in diameter whereas the short, straight filaments were between 4 and 7 nm. Larger diameter filaments (10 nm) were seen more rarely and were always of the long, curving variety. Occasionally, single microtubules coursed smoothly through the dense filamentous meshwork. A fine granular material was tightly packed between the filaments. The diameter of individual granules was extremely variable ranging from 4 to 27 nm with a mean of 15 nm (±0.3 SEM, n = 200). This material was uniformly distributed throughout the cytoplasm but appeared to stain more darkly in some regions where large, dark-staining granules, presumably ribosomes, were concentrated.

Addition of tannic acid to the primary substitution fluid increased the overall density of the cytoplasm although long curving filaments stood out with increased contrast. Hafnium staining darkened the microtubules in relation to filaments and granular material, making it easier to follow microtubules through the cell. Nevertheless, the overall appearance of the cytoplasm remained the same with the various stains.

To determine whether the cell membrane was intact, we shadowed some preparations with a thin layer of nickel/carbon (Fig. 4). The grain of the nickel layer allowed detection of the upper membrane surface while retaining enough transparency to view the underlying cytoplasm. In most well-frozen preparations the membrane was intact, without any detectable holes or fissures.

**Physically Lysed Cells**

Some cells were physically lysed by quickly drawing a piece of filter paper over the grid immediately (2 s) before freezing. Cells were in various degrees of lysis with various degrees of release of their granular components. Some cells were found that were devoid of most of their granular material. The degree or relative length of time of disruption could be roughly monitored by the appearance of mitochondria, which became swollen and then beaded after lysis.

The cytoskeletons of physically lysed cells contained an interconnected meshwork of filaments of various diameters very similar to those of intact cells. Many Y-shaped, short, straight filaments were seen interconnecting longer, thicker filaments (Fig. 5). When cells were lysed under special buffer conditions to preserve labile components (30), there was no detectable difference in the meshwork of filaments, other than a slight increase in the number of microtubules, when compared to those lysed in Steinberg's solution. This was presumably because the time between lysis and freezing was so short. The filamentous meshworks were sufficiently intact to discern

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**Figure 4** Stereo view of a nickel/carbon shadowed cell. The cell was prepared in the same manner as described for the cell shown in Fig. 1. This region is from a small, thin extended process. (inset) The grainy nickel layer allows detection of the upper membrane surface. M, mitochondria; V, vesicle. 1,000 kV. × 26,000. (inset) × 114,000.
different regions of the cell such as the leading edge and other thinly spread areas. Although the cytoskeleton of physically lysed cells appeared to be organized in the same manner as the cytoskeleton of intact cells, the concentration of filaments, which varied somewhat from cell to cell, appeared less.

Detergent-lysed Cells

The detergents Triton X-100 or saponin were used to lyse cells under buffer conditions typically used to maintain cytoskeletal elements (30). Triton extraction was more easily controlled and reproducible than physical lysis, but did not allow observations on interactions between membranes and the cytoskeleton. Saponin was intermediate between Triton extraction and physical lysis in that membranes and organelles were still detectable in some saponin-treated cells. The filamentous meshwork with either detergent contained long, curving filaments interconnected by short straight filaments (Fig. 6) and was similar to that seen with physical lysis. Microtubules were abundant in detergent-lysed cells because of the special buffer conditions. Granular material was absent from detergent-lysed cells in all but a very few instances. Again, there was some variation from cell to cell but the filament meshwork was less concentrated than in intact cells, although the general organization was the same. We could not detect any major differences between these preparations and those obtained by detergent lysis and conventional chemical fixation (30).
Filipin-lysed Cells

Filipin is similar to saponin or digitonin in that it interacts with membrane cholesterol and can lyse cells (5, 27). It differs, however, from saponin or digitonin as it does not act as a surfactant and therefore would not be expected to extract membranes to the same degree as the detergents (19). Stereo images of filipin-treated cells showed that they retained much of their cell membrane, although small holes often appeared and occasionally sheets of collapsed or torn off membrane were seen (Fig. 7). The cytoplasm of filipin-lysed cells was devoid of granular material except for the dark-staining granules that probably represent ribosomes. Membrane-bound organelles were prominent and distributed in a manner similar to that seen in some of the physically lysed cells. Filaments seemed well preserved and appeared to be more numerous than in detergent-lysed cells, though their overall organization was very similar (Fig. 7).

Thin Sections of Directly Frozen, Freeze-substituted Cells

Sections 150-nm thick were cut en face through plastic-embedded, freeze-substituted cells, and stereo micrographs were taken at 120 kV for comparison with micrographs of whole mounts. Prolonged pre-embedding staining was necessary to make cellular components more electron scattering than the resin (38). Thin sections perpendicular to the surface of the grid were also useful in evaluating conclusions about the quality of freezing. Based on the size of ice crystal–induced spaces, we could classify freezing quality into the same categories established for the whole mount images. The thin cross sections also provided measurements of cell thickness. Thinly spread cells such as fibroblast and epithelial cells were between 1.0- and 1.5-μm thick at their center (nuclear region) and 0.1- and 0.06-μm thick in peripheral regions devoid of organelles.

A dense meshwork of filaments was readily apparent in stereo views of en face sections through thin peripheral regions of embedded cells (Fig. 8). This meshwork had the same characteristics and sizes as the meshwork in whole mounts. Less apparent in thin sections was the granular material that is so obvious in whole mounts. However, a few granules of the same diameters (15–18 nm) as those seen in whole mounts were found. Occasionally, the first sections cut parallel to the substrate revealed edges of cells that were thinner than the section and therefore wholly contained within it (see Fig. 10). Comparison of these views with views of whole mounts (e.g., compare Figs. 9 and 10) showed a close correspondence, especially between the relative contribution of granular and filamentous components in the two images.

Fixed, Frozen, Freeze-substituted Cells

Cytoplasmic structure in directly frozen, freeze-substituted cells differed from that in whole mounts of cells prepared by conventional chemical fixation (4, 38). In order to determine whether the initial aqueous glutaraldehyde fixation contributed to these differences, some cells were fixed with glutaraldehyde before freezing to separate effects of initial glutaraldehyde fixation from effects of aqueous osmium fixation or dehydration (33). Freezing artifacts were less frequent in fixed cells than in directly frozen cells possibly because glutaraldehyde has a cryoprotectant effect, and the fixed cytoplasm is less suscep-
FIGURE 8 Stereo view of a thin section through a cell that was directly frozen, freeze-substituted in acetone with a sequential fixative combination of acrolein-tannic acid/osmium/glutaraldehyde, and then embedded in Araldite. The en face section was taken from a peripheral region of the cell. A dense meshwork of cytoplasmic filaments (arrows) is apparent. Dark-staining granules (arrowheads) are occasionally interspersed between the filaments. The graininess of the image results from interactions between the grid stain (lead citrate and uranyl acetate) and the embedding plastic. 120 kV. × 82,000.

Initial glutaraldehyde fixation caused an overall increase in the transparency of the cytoplasm (Fig. 11) because much of the fine granular material was not evident. Filaments therefore stood out against a relatively empty background. Some densely stained granular material was evident, especially after prolonged uranyl acetate staining, but individual granules were much more uniform in diameter and much less frequent than in directly frozen material; these were probably ribosomes, because their size and staining was similar to that of ribosomes in thin sections.

The exact appearance of the fine filamentous meshwork and the ground substance depended on the osmolarity of the buffer used during initial fixation. Working under the assumption that glutaraldehyde can contribute to the total osmolarity of the fixative (14), we fixed some cells using hypotonic buffers (72 mOsmol); the buffer plus glutaraldehyde and tannic acid was approximately isotonic with the Steinberg's solution (134 mOsmol). Cells fixed in this manner had an interconnecting meshwork of distinct filaments of uniform diameter along their length (Fig. 11 A). The filaments were either long and slightly curved, or short and straight. If the buffer was only slightly hypotonic (120 mOsmol), the meshwork was tighter and filaments were much less distinct, more variable in thickness, and more curved (Fig. 11 B). The application of hypertonic buffers (180 mOsmol) resulted in an even tighter, branching network of mostly curved, indistinct filaments of various diameters that were usually thicker at branch points (Fig. 11 C). Close examination of the cytoplasmic meshwork in cells prepared with hypertonic fixatives indicated that the heterogeneity in filament diameter was partially a result of the intertwining or close apposition of two or more filaments. In preparations fixed in hypotonic buffers, the dark-staining granular material (~19 nm diam) was often segregated, producing clear areas between dense granular areas. Segregation of dark-staining granular material was less noticeable in specimens fixed with isotonic buffers.

Filament Junctions

Numerous junctions between filaments were detected in all preparations. However, they were most clearly seen in filipin-treated cells (Fig. 7) and cells fixed with glutaraldehyde in a hypertonic buffer (Fig. 11). The Y-shaped junctions were the most frequent, although right-angles or T-junctions were also seen. Complicated junctions with four to five arms were occasionally found at cell margins (Fig. 5).

Sometimes it was possible to see a contact between two filaments that resembled a Y-junction but on closer inspection appeared to represent the meeting of two filaments which adhered along their length to form a "doublet" (Fig. 6). In addition, it was apparent that two crossing fibers could bend at the point of contact (Fig. 7). Whether it was ever necessary to interpret junctions as actual filament branching is not clear.
Filament Diameters

Many of the filaments in the meshwork of whole mount cells were 7-8 nm diam, regardless of whether the cells were directly frozen, lysed, or fixed before freezing (Fig. 12). Smaller filaments (4-6 nm) were also present in all preparations, although their contribution to the total population varied from 11 to 43% (Fig. 12). This variation could result from loss of some of the smaller, interconnecting filaments in lysed cells. Cells lysed with Triton or saponin before freezing had larger proportions of filaments 9-14 nm diam than did directly frozen or physically lysed cells, apparently from a decoration of smaller filaments with debris (Fig. 6).

Stress fiber filaments that are presumed to represent actin
Figure 11  Stereo views of three different cells fixed with glutaraldehyde in different osmolarity buffers before freezing and freeze-substitution in osmium/acetone. All three grids were freeze-substituted and critical-point-dried in the same run. (A) Cell fixed using a hypotonic buffer (0.05 M HEPES; 72 mOsmol). Total fixative was approximately isotonic with Steinberg’s solution with which cells were equilibrated immediately before fixation. Filaments are distinct, often straight or gently curved, and organized in a manner similar to those in direct frozen or lysed/frozen cells. The appearance of filaments is particularly close to that in filipin-lysed cells (Fig. 7). Single or clumps of dark-staining granules are numerous (arrowheads), but otherwise spaces between filaments are empty. Arrows indicate filament junctions. (B) Cell was fixed using a slightly hypotonic buffer (0.075 mM HEPES; 120 mOsmol). Total fixative was hypertonic. Individual filaments are more difficult to follow for any length because of their increased overlap and curvature. There appears to be more filament junctions than in A. (C) Cell was fixed using an hypertonic buffer (0.1 M HEPES; 180 mOsmol). Individual filaments are very difficult to follow because of the amount of overlap and close lengthwise association. Also, they are curved and make numerous bends. This network of filaments is studded with dark-staining granules. Filament junctions are difficult to see clearly and when detected appear thickened (arrowhead). (A–C) 200 kV. × 74,000.
FIGURE 12  Comparison of filament diameters from cells prepared by direct freezing, by lysis, or by fixation and then freezing. Measurements were made on all clearly defined filaments in high magnification micrographs within a sample area. The 7- and 8-nm bins are darkened. Filaments too large and too small to be actin are present in different proportions after different treatments.
filaments were also measured in intact and lysed cells. Stress fiber filaments had a uniform diameter within a preparation and when results from the different preparations were pooled, they still indicate a fairly homogeneous population of diameters, generally 7–8 nm. (Fig. 12).

Diameters of filaments in thin sections were close to those in whole mounts (Fig. 12), though a smaller proportion of filaments were in the 4–6-nm category. However, the finer filaments were difficult to measure because filament boundaries are not as easy to see in the more grainy images obtained from thin sections. But it is still possible to confirm the contribution of fine filaments to the meshwork by a careful examination of stereo pairs of thin sections (Fig. 8).

DISCUSSION

The cytoplasm in thin peripheral areas of cells prepared as whole mounts by direct-freezing, freeze-substitution, and critical-point drying contains a dense filamentous meshwork embedded in a granular ground substance. A major component of the highly interconnected meshwork are filaments with a uniform diameter of 7–8 nm. These filaments are presumably F-actin because stress fibers in our preparations were composed primarily of similar 7–8-nm filaments that are actin (33); other works on detergent-extracted fixed cells labeled with myosin SI fragments support this interpretation (24, 30). In addition, regions of the meshwork resemble the branching patterns in actin-actin binding protein gels viewed in shadowed, critical-point-dried preparations (18). However, we also find finer filaments, 4–6 nm diam, intercalated in the meshwork which may be related to the fine filaments that do not label with SI in detergent extracted, fixed cells (30).

Recently, 5-nm filaments of the intestinal brush border have been identified as belonging to the spectrin/fodrin family (11). These results together with the present one suggest that spectrin-like filaments could be ubiquitous and supports the interpretation that the 4–6-nm filaments belong to the spectrin/fodrin family, although the presence of filaments made up of myosin monomers or stages of myosin monomer association can not be ruled out.

This view of cytoplasm differs in two important ways from the view derived from examination with high-voltage electron microscopy of fixed whole mounts of cultured cells (4, 23, 24, 38). The first difference is that the filaments that make up the dense meshwork are discrete rods of uniform diameter along their length, and there is no discernible difference, other than the total numbers of filaments, between the structural organization of the meshwork in gently lysed cells and intact cells. A main feature of this organization are the many contacts between the individual discrete components. This aspect of the picture is similar to, if not the same as, that seen in cells extracted with detergents and fixed under appropriate conditions to preserve structural elements (30).

The structural organization of the cytoplasm as we observe it is not adequately characterized by an image of discrete filaments woven into a complex fabric, as has been suggested by examination of detergent-extracted, fixed, frozen, and etched cells (9), nor by the image of curvilinear strands in a continuous trabeculum as visualized on the basis of examining whole mounts of fixed cells with the electron microscope (38). The former picture does not take into account the numerous interconnections between individual filaments and the latter does not describe the discrete nature of the individual structural elements.

A second important difference is that our preparative methods show that the cytoplasm contains a fine granular material packed between filaments that is quickly lost upon cell lysis. This granular material effectively takes up all residual space between filaments and is distributed throughout the cytoplasm. This is apparent in the fine granular material seen in replicas of directly frozen, etched cells (9, 28). The diameters of granules in optimally frozen cells, although variable, are all greater than those of granules seen in optimally frozen albumin solutions. We therefore assume that the granular material as a whole represents a concentrated solution of protein whose individual components are complexes of soluble proteins.

Primary fixation in aqueous aldehyde obscures this granular material, with the exception of the large, dark-staining granules that probably represent ribosomes. This finding suggests one reason why this granular material has not been previously detected in whole mounts of cells (4, 38). However, the granular material was also not detected in whole mounts of direct frozen, freeze-dried, or freeze-substituted PtK2 cells (22). We do not know the reason for this discrepancy, although we can suggest several possibilities. Unfixed and therefore unsupported cytoplasm should be subject to the same artifacts during freeze-drying as it is during deep etching. Extensive etching of directly frozen material is known to give the impression of "clean" filaments and cross-bridges, and the intervention granular material is lost (28). That granular material was not previously seen in freeze-substituted whole mounts may result from the choice of a highly polar solvent, 2-methoxyethanol, for freeze-substitution (22). The more polar solvents cause a dramatic decrease in the amount of detectable granular material which may be due to increased extraction of soluble components by the substitution fluid or may simply reflect differences in staining (Appendix).

Freezing that results in ice crystals >5 nm also produced images of clean, curving, relatively thick filaments without intervening granular material. Similar images resulted from poor freezing of solutions of albumin processed by freeze-substitution and critical-point drying. The formation of ice crystals concentrates soluble materials into boundary layers between crystals that give the impression of irregular filament-like structures bordering open spaces in the cytoplasm or protein solution (20).

The only difference between images of whole mounts of cells and thin-sectioned cells is accounted for by differences in specimen thickness and contrast. The decreased contrast in thin sections makes the lightly stained granular material more difficult to see. However, the organization of filaments seems to be the same with these two techniques, though the filamentous meshworks in directly frozen cells, whether viewed in whole mounts or thin sections, appear more distinct and uniformly spaced than in fixed cells. A major reason for this difference seems to be the osmotic effects of glutaraldehyde (14), which, like glycerol, penetrates the cell slowly (36). If fixation occurs before equilibration, the cell will be fixed in a dehydrated state and will appear shrunken. This may be the explanation for why cells initially fixed in hypotonic buffer-fixative solutions have filamentous meshworks that are most similar to directly frozen cells.

Fixation of cultured cells in a saponin, tannic acid, and glutaraldehyde solution followed by a low concentration of osmium (16) supports our interpretation of the effects of fixation on cytoplasmic structure. By permeabilizing the cells...
with saponin, osmotic effects are circumvented and it is possible that filaments are fixed in a relatively undistorted position. This could also be the explanation for why there have been such differences between the appearance of filamentous meshworks in whole mount cells fixed while intact as compared with those fixed after extraction with the detergent Triton X-100 (15, 24, 30). Clearly, cells can be initially fixed under appropriate conditions to give an accurate picture of the organization of their cytoplasmic filaments.

APPENDIX

Freeze-substitution

Mixtures of osmium and acetone are typically used for freeze-substitution of tissue to be thin-sectioned (37). However, we soon found that typical osmium-acetone combinations were inadequate for directly frozen cells because of their tendency to shrink during the critical-point drying. Osmium/acetone freeze-substitution is also known to be an inadequate preparation for scanning electron microscopy (1). Shrinkage could be partially alleviated by using high concentrations of osmium (3-5%) and warming the specimens to room temperature while in the osmium/acetone. However, the dense staining that resulted obscured most of the details in cell whole mounts. Another concern was that osmium could have detrimental effects on actin fibers similar to those occurring in aqueous osmium solutions (17). We therefore decided to look for a better method of freeze-substituting specimens to be viewed as whole mounts. Several subjective criteria were used to determine the best methods: (a) the state of preservation of readily recognizable fibrous elements, such as microtubules and stress fibers; (b) the state of preservation of organelles such as mitochondria and endoplasmic reticulum; (c) the continuity of cell membranes; (d) the preservation of fine cell projections such as filopodia; and (e) the relative density and appearance of the cytoplasmic ground substance. To insure that artifacts induced by freeze-substitution were clearly distinguishable from freezing artifacts, we subjected at least six grids to each variation of the substitution procedure and then scanned them in the electron microscope. Recrystallization of ice (7) rarely occurred as long as high-grade solvents free of water were used. Recrystallization was distinguished from poor initial freezing by the ubiquity of large ice crystals in all the samples from a run. Recrystallization occurs when samples are warmed before substitution is complete (35).

Direct frozen whole mounts were subjected to over 60 different combinations of substitution solutions, temperatures, and times. In the most successful method, whole mounts were substituted in 10% acrolein in acetone usually with 0.2% tannic acid at −80°C for 10-18 h, warmed to −55°C, and transferred (after a rinse in acetone) to 0.2% osmium tetroxide in acetone that was then warmed to −20°C. After 2-3 h at −20°C the whole mounts were rinsed, transferred to anhydrous 10% glutaraldehyde in a methanol/acetone mixture (40%/50%), warmed to 0°C, and maintained at that temperature overnight. Specimens were then washed with methanol (40 min) and stained with 0.1-0.5% uranyl acetate in methanol or acetone (30-45 min at 0°C) followed by (0.5-1.0%) hafnium chloride in methanol or acetone for 20-45 min at room temperature.

When whole mounts were to be embedded and then thin-sectioned, the procedure was modified to increase the staining and fixation. The osmium concentration was increased to 0.4% and the time at −20°C in osmium was increased from 2 to 8 h. In addition, the length of staining with uranyl acetate was increased to ~12 h and with hafnium chloride to 2 h. The results of the trials with different solvents, fixatives, and stains, which provide a rationale for this method, are detailed below.

Solvents

Solvents or solvent combinations tried for freeze-substitution were tetrahydrofuran, acetone, ethanol, 90% ethanol/10% H2O, 90% ethanol/10% glycerol, methanol, and 60% methanol/40% ethylene glycol (1).

The more polar solvents such as methanol and ethanol caused partial loss of cell membranes and some shrinkage and partial extraction of the cytoplasm even in the presence of the best fixative combinations (see below). The extraction of cytoplasmic components was indicated by the decreased density of the cytoplasm compared with samples substituted in acetone or tetrahydrofuran. Addition of water, glycerol, or ethylene glycol to slow the rate of substitution prevented shrinkage but had no noticeable effects on preserving membranes or preventing extraction of cytoplasmic components.

No shrinkage could be detected when either acetone or tetrahydrofuran was used with the best fixatives. Cell membranes remained intact in these solvents as long as substitution times at −80°C were <48 h. Substitution times >64 h caused loss of membranes, as did warming to >−40°C without osmium fixation.

Fixatives and Stains

Substantial chemical fixation during freeze-substitution was necessary to prevent shrinkage induced by critical-point drying. Osmium fixation alone was not adequate because concentrations, times, and temperature of osmium fixation that gave well-stained preparations did not prevent shrinkage. For this reason, we tried various combinations of fixatives.

Acrolein (10%) was the most effective fixative at cold temperatures and was the only fixative to give indications of fixing (preserved shrinkage) at −80°C. However, it is difficult to give an accurate indication of the time necessary for fixation at −80°C because the time needed for substitution is unknown. The minimum time that was tried and found to be effective for substitution with 10% acrolein in acetone was 10 h. Although acrolein provided good structural preservation of the cytoplasm, it gave minimal fixation of membranes, even at temperatures of 0°C.

Osmium even at concentrations of 5%, had no apparent effect at the substitution temperature (~−80°C) (37). The lowest temperature at which osmium showed signs of fixing (preserved structure and stained membranes) was between −40°C and −50°C. By independently adjusting concentration, time, and temperature of exposure to osmium we came up with an exposure just sufficient for preservation and staining of membranes, in order to avoid possible detrimental effects on actin filaments. The useful concentrations were between 0.2 and 0.4% at temperatures up to −20°C and times ranging from 6 to 14 h.

Glutaraldehyde (10%) was the least effective fixative at cold temperatures. No fixation was observed at temperatures −<−20°C (39), and from −20°C to +4°C, fixation that helped prevent shrinkage occurred after fixation at colder temperatures by acrolein or osmium.

Tannic acid is not generally considered a fixative, but it is known to protect actin filaments from fragmentation by aqueous osmium (16). It also stains actin filaments and membranes directly, presumably by increasing the reduction of osmium (16). Tannic acid was useful in freeze-substitution both as a stain and as a preserving agent. Addition of 0.2% tannic acid (desiccated for 4 h at 80°C) to the primary substitution fluid increased subsequent osmium staining of membranes, ground substance, and filaments. Membranes were also less likely to show the discontinuities that sometimes appeared when lower concentrations of osmium alone were used. Tannic acid was also an effective general stain when applied at high concentrations and warmer temperatures after osmium fixation.

Uranyl acetate stabilizes membranes following aldehyde fixation (31, 32), and we found that it could help stabilize membranes at low temperatures depending on the solvent. Membranes were extracted in the more polar solvents but were generally intact when substituted in ethanol or acetone. Uranyl acetate increased the contrast of membranes and filaments after fixation by aldehydes and osmium. Its staining properties were not dependent simply upon concentration, time, and temperature but also on the solvent. For instance, 0.5% uranyl acetate in acetone applied for 30 min at 0°C gave more intense staining than the same concentration in methanol applied for the same time and at the same temperature. Hafnium chloride in methanol was especially useful for impregnating contrast to microtubules when used in combination with uranyl acetate. Like uranyl acetate, it gave more intense, general staining when used in acetone than when used in more polar solvents such as methanol.

Conventional fixation by aqueous solution of fixatives produces the best results when used in combinations; no fixative has been found that will adequately fix all cellular constituents. The most widely used combination is aldehyde fixation followed by postfixation in osmium tetroxide. The best and most consistently reproducible fixation during the freeze-substitution process is a similar combination of fixatives. However, the constraints imposed by the freeze-substitution temperatures on the activity of the various fixatives partially dictate the particular combination.

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