Immunolocalization of Actin in *Paramecium* Cells

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**SUMMARY**

We have selected a conserved immunogenic region from several actin genes of *Paramecium*, recently cloned in our laboratory, to prepare antibodies for Western blots and immunolocalization. According to cell fractionation analysis, most actin is structure-bound. Immunofluorescence shows signal enriched in the cell cortex, notably around ciliary basal bodies (identified by anti-centrin antibodies), as well as around the oral cavity, at the cytoproct and in association with vacuoles (phagosomes) up to several \( \mu \text{m} \) in size. Subtle strands run throughout the cell body. Postembedding immunogold labeling/EM analysis shows that actin in the cell cortex emanates, together with the infraciliary lattice, from basal bodies to around trichocyst tips. Label was also enriched around vacuoles and vesicles of different size including “discoidal” vesicles that serve the formation of new phagosomes. By all methods used, we show actin in cilia. Although none of the structurally well-defined filament systems in *Paramecium* are exclusively formed by actin, actin does display some ordered, though not very conspicuous, arrays throughout the cell. F-actin may somehow serve vesicle trafficking and as a cytoplasmic scaffold. This is particularly supported by the postembedding/EM labeling analysis we used, which would hardly allow for any large-scale redistribution during preparation. *(J Histochem Cytochem 52:1543–1559, 2004)*

**KEY WORDS**

actin
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*Paramecium*


Different actin isoforms occurring in many organisms may serve specific functions in the respective cells (Pollard et al. 2000; Wagner et al. 2002). For localization, antibodies (ABs) may be used at the light microscope (LM) and electron microscope (EM) levels, as well as for Western blots. Bicyclic peptide toxins, phalloidin or jasplakinolide, can bind rather specifically to F-actin, thus allowing fluorescence labeling (Wieland and Faulstich 1978; Bubb et al. 2000). This or the alternative approach, F-actin disruption by toxins of the
type cytochalasin B and D or latrunculin A, is also widely used for functional analyses also with ciliates (see below).

In previous times, mainly before molecular biology approaches could be undertaken, biochemical, functional, and immunolocalization studies were tried to probe the potential function of F-actin in ciliates such as *Paramecium* (Tiggesmann and Plattner 1981; Cohen et al. 1984; Fok et al. 1985; Kersken et al. 1986a,b), *Tetrahymena* (Mitchell and Zimmerman 1985; Hirno et al. 1987b, 1989; Hoey and Gavin 1992), *Pseudomicrotborax* (Hauser et al. 1980), *Histrichulus* (Pérez-Romero et al. 1999), *Climacostomum* (Fahrni 1992), and *Spirostomum* (Zackoff and Hufnagel 1998). However, with ciliates, F-actin–disrupting drugs frequently had to be used in conspicuously high concentrations to abolish, e.g., phagocytosis (Fok et al. 1987; Zackroff and Hufnagel 1998, 2002). With a variety of protozoa of the phylum Alveolata, actin genes or partial sequences of it have been cloned. This holds in particular to abolishes, e.g., phagocytosis (Fok et al. 1987; Zackroff and Hufnagel 1998, 2002). With a variety of protozoa of the phylum Alveolata, actin genes or partial sequences of it have been cloned. This holds in particular for ciliates, such as *Tetrahymena* (Zimmerman et al. 1983; Cupples and Pearlman 1986; Hirno et al. 1987a) and *Paramecium* (Díaz-Ramos et al. 1998), but also for their pathogenic relatives of the group of Apicomplexa such as *Toxoplasma* (Delbac et al. 2001).

Our present analysis also addresses some special subcellular structures in *Paramecium* cells that contain multiple filament systems (Allen 1971; Cohen et al. 1984, 1987; Cohen and Beisson 1988; Keryer et al. 1990a,b; Allen et al. 1998; Beisson et al. 2001; Clérot et al. 2001). We focus on regions with dense-core secretory vesicles (“trichocysts”), cortical filament bundles (“infraciliary lattice,” cf. Allen 1971, 1988), the narrow space between the plasma membrane and tightly attached cortical Ca$^{2+}$-stores (“alveolar sacs,” see Plattner and Klauke 2001), in addition to abundant vesicles of the phagolysosomal and recycling system (Fok and Allen 1990; Allen and Fok 2000). Recent cloning of several actin genes of *Paramecium tetraurelia* in our laboratory opened up a new way to structural localization with this cell, whose regular “design” facilitates such studies. So far, studies on actin in *Paramecium* have not addressed all relevant aspects, and many aspects have remained controversial.

**Materials and Methods**

**Stocks and Cultures**

The wild-type strain of *P. tetraurelia* used was stock 7S. Cells were cultivated in a decoction of dried lettuce monoxenically inoculated with *Enterobacter aerogenes* as a food organism, supplemented with 0.4 μg·ml$^{-1}$ β-sitosterol (Sonnewborn 1970). For subcellular fractionation, we used axenic cultures (Kaneshiro et al. 1979). Cells were grown at 25°C to early stationary phase as previously described (Kissmehl et al. 1996).

**Expression of Paramecium Actin-specific Peptides in Escherichia coli**

For heterologous expression of actin-specific peptides we selected the amino acid sequence of actin1-1 [accession number AJ537442](https://www.ncbi.nlm.nih.gov/nuccore/AJ537442). After changing all deviant *Paramecium* glutamine codons (TAA and TAG) into universal glutamine codons (CAA and CAG) by PCR methods, the coding regions of either E57-P243 (N-terminal region) or L251-G366 (C-terminal region) of *Paramecium* actin1-1 were cloned into the Xhol/BamHI restriction sites of pET 16b expression vector of the pET System from Novagen (Madison, WI) which employs a His$_{10}$ tag for purification of the recombinant peptides.

**Purification of Recombinant Actin1-1 Peptides**

Recombinant actin1-1 peptides, actin1-1.E57-P243 and actin1-1.L251-G366 were purified by affinity chromatography on Ni$^{2+}$-nitrilotriacetate agarose under native conditions, as recommended by the manufacturer (Novagen). The recombinant peptides were eluted with a step gradient, 100 to 1000 mM imidazole in 50 mM sodium phosphate (pH 6.0) with 300 mM NaCl added. The fractions collected were analyzed on SDS polyacrylamide gels, and those containing the recombinant peptides were pooled and dialyzed in phosphate-buffered saline (PBS).

**Antibodies Used**

Abs against the two recombinant actin peptides, actin1-1.E57-P243 and actin1-1.L251-G366 were raised either in rabbits or mice. After several boosts, positive sera were taken at day 60 and purified by two subsequent chromatography steps, a first step on a His-tag peptide column (24-amino acid peptide, to remove His tag-specific ABs), followed by an affinity step on the corresponding actin1-1 peptide. One of these ABs recognizes the N-terminal and the other the C-terminal region of actin1-1, yet results achieved in this study were indistinguishable with either type of ABs. Therefore, no further distinction is made, unless indicated. We used the sequence of *Paramecium* actin1-1 because it is rather similar for numerous isoforms that we have cloned (R. Kissmehl, J. Mansfeld, E. Wagner, I. Sehring, H. Plattner, unpublished data) and thus should allow us to establish an overall distribution of actin, notably of F-actin, in *Paramecium*.

Mouse polyclonal ABs against *Paramecium* actin1-1 were selectively used for the colocalization at the LM level, in conjunction with an anti-centrin (*Dictyostelium discoideum*) polyclonal AB produced in rabbits (designation HisDd-Centrin2 from R. Gräf, University of Munich) used to identify ciliary basal bodies (Daunderer et al. 2001).

**Cell Fractionation**

Cells were deciliated by a Mn$^{2+}$-shock (for details, see below) and cilia were purified by differential centrifugation (Nelson 1995). Whole-cell homogenates were prepared in phase buffer (20 mM Tris-maleate, 20 mM NaOH, 20 mM NaCl, 250 mM sucrose, pH 7.0) by ~100 hand strokes in a glass homogenizer equipped with a Teflon pestle. Soluble and particulate fractions were separated by centrifugation at 100,000 × g for 60 min at 4°C. Cell surface complexes (“cortex”) were prepared according to Lumpert et al. (1990),
and trichocysts were isolated by the method of Glas-Albrecht and Plattner (1990). A protease-inhibitor cocktail containing 15 μM pepstatin, 100 μM aprotinin, 100 μM leupeptin, 0.26 mM TAME, 28 μM E64, and 0.2 mM Pefabloc SC was used throughout.

**Electrophoretic Techniques and Western Blot Analysis**

Protein samples were denatured by boiling for 3 min in sample buffer (0.4 M Tris–HCl, 1% SDS, 0.5% DTT, 20% glycerol, pH 8.0) and subjected to electrophoresis on linear gradient (5–20%) SDS polyacrylamide gels using the discontinuous buffer system of Laemmli (1970). Before electrophoresis, samples were alkylated for 30 min at 20°C by 2% iodoacetamide. Protein standards were used in accordance with manufacturer directions. Gels were either stained with Coomassie blue R250 or prepared for electrophoretic protein transfer onto nitrocellulose membranes. Protein blotting was performed at 2 mA/cm² for 1 hr according to the technique of Kyhse-Andersen (1989) using the semidyed blotter from BioRad (Munich, Germany). ABs were diluted 1:100 with manufacturer directions. Gels were either stained with preadsorbed with the original antigen.

**Immunofluorescence Labeling**

**Basic Procedure.** Cells were washed twice in 5 mM Pipes buffer, pH 7.0, containing 1 mM KCl and 1 mM CaCl₂. Cells were fixed in 4% (w/v) freshly depolymerized formaldehyde with 0.5% digitonin, 1 mM ATP, 10 mM MgCl₂, and 10 mM KCl added, further fixed for 1 hr according to the technique of Taufkirchen (1970). Before electrophoresis, samples were alkylated for 30 min at 20°C by 2% iodoacetamide. Protein standards were used in accordance with manufacturer directions. Gels were either stained with Coomassie blue R250 or prepared for electrophoretic protein transfer onto nitrocellulose membranes. Protein blotting was performed at 2 mA/cm² for 1 hr according to the technique of Kyhse-Andersen (1989) using the semidyed blotter from BioRad (Munich, Germany). ABs were diluted 1:100 with manufacturer directions. Gels were either stained with preadsorbed with the original antigen.

**Deciliated Cells.** Cells were washed twice in 5 mM Pipes buffer, pH 7.0, containing 1 mM KCl and CaCl₂, at room temperature and suspended in 50 mM MnCl₂ solution in 10 mM Tris-HCl, pH 7.2. After 2 min at 4°C, cells were removed by centrifugation and resuspended in the same solution. After 10 min of gentle shaking, 90–95% of cells were deciliated. Deciliated cells were removed by centrifugation and washed twice in Pipes buffer before further use.

**Fixation and Embedding for Postembedding EM Analysis**

Using a quenched-flow apparatus (Knoll et al. 1991), *Paramecium* cells were rapidly injected into 8% formaldehyde plus 0.1% glutaraldehyde dissolved in Pipes buffer, pH 7.2 (0C), with 1 mM KCl and CaCl₂ each added, further fixed for 60 min at 4C, washed in PBS (pH 7.4) + 50 mM glycine (2 × 10 min), dehydrated by increasing ethanol concentrations (30%, 50%, 70%, 90%, 96%, 2 × 10 min each, and 2 × 100%, 30 min each), and impregnated with LR Gold resin (London Resin, London, UK) at 0°C, with two changes in 2-hr intervals each and then overnight, followed by UV-light polymerization at −35°C for 72 hr.

**Immunogold Labeling and EM Analysis**

**Postembedding Method.** Ultrathin sections mounted on formvar-coated Ni grids were pretreated (2 × 10 min) with 20 μl of PBS, then for 10 min with PBS with 50 mM glycine added, and finally immersed in PBS supplemented with 0.5% BSA and 0.5% goat serum (2 × 10 min, room temperature), to eliminate nonspecific gold adsorption. Grids were then incubated with rabbit AB, diluted 1:20 in PBS supplemented with 0.3% BSA-c (BioTrend, Köln, Germany), pH 7.4, 1 hr at room temperature. BSA-c as an acetylated form reduces nonspecific adsorption of gold conjugates due to increased net charge.

**Preembedding Labeling.** Without exception, cells were fixed with 8% formaldehyde + 0.1% glutaraldehyde and simultaneously treated with digitonin (Sigma) and the other additives, as described above for LM analysis of deciliated cells, incubated with primary rabbit ABs against *Paramecium* actin1-1, followed by Au₅-conjugated second ABs, with the aim to make the narrow subplasmalemmal space accessible. After embedding in LR Gold (London Resin), sections were additionally subjected to the postembedding labeling procedure with the same primary and secondary ABs, respectively.
Further Processing and Quantitative Evaluation. After labeling, sections were rinsed with distilled water, fixed for 5 min with 2% glutaraldehyde, and routinely stained for 3 min with 2% aqueous uranyl acetate (unbuffered, pH ~4.5). EM micrographs were taken at defined magnifications and enlarged to ×77,000. Au5 grains were counted and referred to area size determined by superposition of square lattices with 5, 10.0 and 20.0 mm spacing, respectively, depending on the size of the structure to be analyzed (Plattner and Zingsheim 1983). The actual area sizes to which the numbers of gold grains were referred were determined from the number of hit points.

Results

Actin-specific ABs, Cell Fractionation, and Western Blot Analysis

Molecular cloning from a pilot sequencing project (Des- sen et al. 2001; Sperling et al. 2002) as well as from the ongoing Paramecium genome project of the Groupe- ment de Recherche Européen at the Genoscope (Evry, France) revealed that P. tetraurelia contains an actin multigene family with at least 30 members, all encoding actin and actin-related proteins with calculated molecular masses ranging between 38 and 45 kD (R. Kissmehl, J. Mansfeld, I. Sehring, E. Wagner, H. Platt- ner, unpublished data). One of them, actin 1-1 (accession number AJ537442), a member of the actin-1 subfamily with rather conserved immunogenic regions (Figure 1), was chosen for heterologous expression in E. coli (after changing all deviant Paramecium glutamin codons into universal glutamine codons) and subsequent production of polyclonal ABs. Various polyclonal ABs were raised against the N-terminal (E57-P243) or C-terminal region (L251-G366, Figure 1), all readily recognizing the recombinant peptides used for immunization when tested in slot blots and Western blots (data not shown). After affinity purification, the actin-specific ABs were further characterized in ELISA and Western blots. Results obtained were similar, whether ABs were used against the N-terminal or the carboxy-terminal region of actin 1-1, confirming their high specificity against actin or actin-specific peptides (data not shown). The following analyses, including Western blots, and LM and EM analyses, have been performed predomin-antly with ABs against the C-terminal region of Paramecium actin1-1 (Figure 1).

Western blots from homogenates display a strong band of 43 kD and a weak one of 40 kD (Figure 2). Such bands also occur in the 100,000 × g pellet, while the 43 kD band is much weaker in the 100,000 × g supernatant. The 43 kD band is typical of actin, while the 40 kD band may represent one of the shorter iso- forms of the actin or actin-related gene products of Paramecium (Kissmehl et al., unpublished data). A 43-kD band also clearly occurs in cilia and in cortices, while it is hardly discernible in the trichocyst fraction. Both the 100,000 × g supernatant and pellet also display some very weak bands of lower size, possibly

Figure 1  Multiple alignment of the C-terminal region of Paramecium actin1-1. Actin-specific sequences from Paramecium tetraurelia (AJ537442), Toxoplasma gondii (P53476), Dicyostelium discoideum (AA052255), Caenorhabditis elegans (X16797), Drosophila melanogaster (NP_523625), Mus musculus (NP_033739), and Homo sapiens (AAH16045) were aligned using the CLUSTALW program. Identical residues are shaded (black), while lesser conserved positions are labeled greyish.
generated by partial proteolysis during fractionation. None of the bands were visible when Western blots were produced with the corresponding preimmune sera or in controls with the second AB alone (data not shown).

Immunofluorescence Labeling

To account for some variability in the immunofluorescence staining, we present typical extremes of CLSM images from double labeling experiments (Figures 3A–3D), with mouse anti-actin FITC-ABs and rabbit anti-centrin Texas Red-ABs, the latter specific for the centrosome in Dictostelium (Dauneder et al. 2001) and basal bodies in Paramecium. This is in contrast to the pattern obtained by the monoclonal AB 20H5 against centrin from Chlamydomonas (Sanders and Salisbury 1994) which in Paramecium brilliantly stains not only basal bodies but also the infraciliary lattice (Klotz et al. 1997; Beisson et al. 2001). Labeling with both anti-actin and anti-centrin ABs in part coincides with ciliary basal bodies of the outer cell surface and along the oral cavity, the outline of the oral cavity, and on the cytoproct. This structure is identified by its “posteroventral” position, size, and shape (Allen 1988). The degree of coincidence (yellow) on basal bodies and in the oral cavity may vary; e.g., it is higher in Figures 3A and 3B than in Figures 3C and 3D. The gradient of coincidence in Figure 3A indicates some differential positioning of the respective antigens along the z-axis.

Figure 3B shows the occurrence of actin around vesicles and vacuoles of various sizes, whereas the position of the red-labeled structures may suggest coincidence with elements of the osmoregulatory system—aspects that have not been followed in any more detail. Figure 3D documents more clearly a cortical actin layer and actin filaments throughout the cytoplasm, frequently in a radial arrangement, and sometimes with local concentration.

We used conventional LM analysis to analyze immuno-FITC labeling of cilia with anti-actin ABs (Figure 4), thus taking advantage of a thicker optical section layer. While intracellular details are largely blurred, ciliary basal bodies and cilia on the outer cell surface are clearly labeled. This may also apply to cilia in the oral cavity, although this is not resolved in Figure 4.

Comparative Analysis of CLSM and Immunogold EM Labeling

For most results achieved by CLSM analysis, we find equivalents in the immunogold EM analyses (Figures 5 to 11), as specified below and summarized in Table 1. Off-cell background is low [2.15 gold grains per μm² ± 0.85 (SEM)], as it is on irrelevant structures, such as mitochondria, trichocyst contents, and alveolar sacs (2.2, 1.4, and 0.3 gold grains per μm², respectively).

After postembedding labeling, gold granules are scattered, yet with specific concentration zones over the cytosolic compartment. This holds for the cell cortex (Figures 6 to 8) with its ciliary basal bodies, as well as for regions adjacent to the oral cavity, including a zone enriched in ciliary basal bodies (Figure 10A) and a zone enriched in recycling vesicles (discoidal vesicles) dedicated to phagosome formation (Figure 10B). It also holds for regions deeper inside the cytoplasm where elements of vesicle trafficking are enriched (Figure 11). Cilia are also labeled at the EM level (Figures 5 and 10, Table 1), just as with the other methods used (Figures 2 and 4). In sum, there is good agreement between LM and EM labeling. Because the cytoproct shows up rarely, we were unable to analyze it at the EM level.

Figure 9 represents experiments with digitonin-permeabilized cells, showing AB-gold labeling in the narrow subplasmalemmal space between the plasmalemma
and the outer side of alveolar sacs, while there is only spurious label occasionally seen after mere section labeling (Figures 5 and 7). Apart from this aspect, little label only is seen in the cell cortex with permeabilized cells (Figure 9). While digitonin permeabilization may be more appropriate than section labeling to show the presence of some actin in the very narrow outermost cytosolic space, particularly when enhanced by additional postembedding labeling (Figure 9), it may cause a serious overall loss of antigen. The abundance of

**Figure 3** Colocalization of actin and centrin (yellow) by CLSM using mouse ABs against actin (green) and rabbit ABs against centrin (red). Two deciliated cells (A,B vs C,D) showing extreme situations of labeling are presented. (A,C) are superficial sections; (B,D) are median focal planes. Note colocalization at basal bodies in top-most focal planes (arrowheads), on the cytoproct (cp) and in parts of the oral cavity (oc). Basal bodies located in layers outside the optical section are preferably red (A) or green (C), thus suggesting a layered arrangement of actin and centrin in these regions of the cell. Note occurrence of actin in the outermost cortex layer particularly in (D, arrowheads) as well as of interior actin clusters probably associated with vacuoles (v in B) and as filament bundles indicated by arrows (D). (B) displays centrin staining at two conspicuous sites where the osmoregulatory system is located (asterisks) and actin labeling associated with large vacuoles (v). Bars = 10 μm.

**Figure 4** Conventional anti-actin AB-fluorescence image of a cell permeabilized under conditions preserving cilia. (A) superficial, and (B) median plane. Note labeling of cilia (ci) in (A) and of their basal bodies (bb) in (A,B), of specks and strands in (B), and of the presumable oral cavity (oc) in (A,B). Bars = 10 μm.
Actin in Paramecium

1549

cortical label after postembedding labeling justifies reliance in this study mainly on the postembedding procedure for further evaluation. Concomitantly, all figures presented with the exception of Figure 9 were obtained by this method.

Specification of Results Obtained with Postembedding Labeling

Beyond the general labeling of the cytosolic compartment of the cell cortex (Figures 5 to 8), we recognize that gold granules are enriched to a variable extent in a variety of structures.

The cytoplasm of cell surface ridges, typical of ciliated protozoa, are labeled (Figures 5 and 6). This also holds for the cytoplasm surrounding the tips of the elongate trichocyst organelles, as shown in cross-section (Figures 5 and 6) and in longitudinal section (Figure 7). The gold label associated with cortical basal bodies is somewhat variable and may in part sit inside this structure, as shown particularly in Figure 8B, where it shows up below the basal plate (Figure 8A). Gold label also occurs adjacent to cortical basal bodies, e.g., in the filamentous mass in Figure 6. This material is associated with the origin of a kinodesmal fiber emanating from a basal body from where the infraciliary lattice also emanates. From there, these filament bundles pass near adjacent trichocyst tips (Figure 5), as established by Allen (1971, 1988). Although the bulk of the latter filament system is made of centrin (Beisson et al. 2001), some actin clearly appears to be associated with it. Gold label also surrounds ghosts from discharged trichocysts (Figure 6).

Table 1 summarizes labeling densities on a quantitative level (gold grains per \( \mu m^2 \)). These are, in decreasing magnitude, as follows: 301.0 Au/\( \mu m^2 \) for cytoplasmic regions around oral cavity and around food vacuoles, 141.5 for cell surface ridges, 111.9 for immediate surroundings of trichocysts, between 89.5 and 95.6 for infraciliary lattice, ciliary basal bodies, and cilia, followed by cortical cytoplasm (37.8) and the complex formed by the plasma membrane and the outer alveolar sacs membrane (25.9 Au/\( \mu m^2 \)). For statistics, see Table 1.

While the abundant filament bundles located in the cytoplasm around the oral cavity are made of materials other than actin (see “Discussion”), the distinct labeling in between such bundles (Figure 10A) again indicates association with actin. As in the cell cortex, some label may be associated with ciliary basal bodies around the oral cavity. Furthermore, we find intense labeling of cytosolic regions enriched in vesicles accumulated near the cytopharynx (Figure 10B). Many are oblong and thus represent discoidal vesicles known to serve membrane recycling from the cytoproct, i.e., formation of new phagocytic vacuoles (Fok and Allen 1988; Allen and Fok 2000). In these domains of the cell, less labeling is seen immediately below the cell membrane than between the adjacent round and discoidal vesicles.

Deeper inside the cell, small vesicles of different diameters are embedded in considerably labeled cytosol, frequently in close association with a large vacuole (Figures 11A and 11B). This arrangement suggests their identity either as lysosomes or as acidosomes in typical arrangement with phagosomes. These interpretations are suggested by the work of Allen and Fok (2000); e.g., considering the flat shape of the large vacuole indicating an early biogenetic stage of a food vacuole. Figure 11B shows association of actin label with parallel microtubular aggregates, the gold label unilaterally concentrated at sites where microtubules enter the section plane. Also in Figure 11B, a heavily labeled “trail” is in direct extension of the adjacent microtubular bundle. This indicates involvement of actin in phago-lysosomal vesicle trafficking, although after the preparation protocol required for immunogold analysis, distinct filaments are difficult to recognize. However, some of these gold aggregates may be the equivalent of the fluorescent strands visualized by anti-actin ABs in Figure 3.

Discussion

Background from Previous Work

Occurrence of most actin in Paramecium in structure-bound form contrasts with the abundance of monomeric actin in Apicomplexa (Sibley 2004), including Toxoplasma (Poupel et al. 2000; Wetzel et al. 2003). This makes fluorescence labeling studies with F-actin–specific drugs feasible. In Paramecium, phalloidin, heavy meromyosin, and DNaseI have clearly revealed labeling of the cell cortex, particularly of ciliary basal bodies (Tiggemann and Plattner 1981; Kersken et al. 1986a,b). Phalloidin also has labeled the nascent food vacuole (Kersken et al. 1986a,b). Concomitantly, cytochalasin B has been reported to inhibit formation of phagocytic vacuoles (Allen and Fok 1983,1985; Fok and Allen 1988). It also inhibits docking of trichocysts (Beisson and Rossignol 1975), and it even can detach docked trichocysts from the cell surface (Pape and Plattner 1990). When phagocytosis has been analyzed with different F-actin–disrupting drugs and analogs, respectively, the requirement of concentrations well above those used with mammalian cells has been confirmed (Beisson and Rossignol 1975; Pape and Plattner 1990; Zackroff and Hufnagel 1998). This is in line with the low sensitivity of F-actin in other ciliates. In total, these data are all compatible with our current results obtained with ABs against the original Paramecium antigen.

Previous attempts to localize actin in Paramecium have led to controversies. One discrepancy concerned
the composition of cortical filament bundles, notably of the infraciliary lattice emanating from ciliary basal bodies. While the bulk of this filament system has been established as centrin (Beisson et al. 2001), this does not necessarily preclude association of centrin filaments with actin, as we can show. Recall that widely different affinity stains for actin, including heavy meromyosin, have resulted in cortical labeling in *Paramecium* (Tiggemann and Plattner 1981; Kersken et al. 1986a,b), as well as in *Tetrahymena* (Méténier 1984). Theoretically, previous LM and preembedding-EM localization studies could have faced the problem of sol-
urable antigen relocation and even loss during permeabilization. This would not easily be possible with the postembedding immuno-EM labeling procedure used now. Another hint to real cortical F-actin localization in *Paramecium* came from the in vivo labeling by injection of fluorescent phalloidin (Kersken et al. 1986a,b), resulting first in cortical labeling and, over longer time periods, in disappearance from the cortex and re-assembly as thick trans-cellular filament bundles of a type not previously seen. Conversely, aberrant phalloidin binding by F-actin formed by some isoforms may preclude labeling (Hirono et al. 1989), while such forms may bind actin-specific ABs.

**Additional Functional Aspects Derived from This Study**

Cortical F-actin is generally required for cyclosis—an actomyosin-based process (Shimmen and Yokota 2004). This is a permanent ongoing process also in *Paramecium* (Sikora et al. 1979), where it serves the delivery of trichocysts to the cell cortex (Aufderheide 1977) and the cycling of phago-lysosomal elements through the cell body (Fok and Allen 1988,1990; Allen and Fok 2000). Myosins occur in *Paramecium* (Cohen et al. 1987), just as in other protists (Gavin 2001).

Our present EM analysis verifies that in the *Paramecium* cell cortex, actin is enriched at ciliary basal bodies, as discussed above on the LM level. From there it emanates to the infraciliary lattice and around trichocyst docking sites. The association of actin with ciliary basal bodies has led to the description of the “basal body cage,” particularly in *Tetrahymena* (Hoey and Gavin 1992), where association with myosin has been demonstrated (Garcés et al. 1995). The loose arrangement of gold label within and around basal bodies, as we see it here, suggests that during permeabilization for LM analysis, F-actin emanating from basal bodies may collapse to a compact arrangement. In

![Figure 7](image-url)
sum, a more loosely arranged cortical F-actin in conjunction with myosin may underlie cytoplasmic streaming and possibly trichocyst docking. Concomitantly, inhibition of trichocyst docking by cytochala-sin B (Beisson and Rossignol 1975) would be compatible with both actin-based transport by cyclosis and enrichment of actin around trichocyst tips (this study).

Figure 8  Postembedding immunogold labeling of ciliary basal bodies (bb) located on the outer cell surface, in longitudinal (A) and in cross-section (B), with additional label on diffuse materials surrounding the basal body (framed in A). Note again absence of label on irrelevant structures, such as alveolar sacs (as), mitochondria (m) and a trichocyst body (tb). Bars = 0.1 \mu m.

Figure 9  Combination of pre- and postembedding immunolabeling shows label in the narrow subplasmalemmal space (at/between arrowheads) between the plasma membrane (pm) and the outer alveolar sacs (as) membrane (oam), with little background on irrelevant structures outside the cytosolic compartment. Note deformation of the cell surface membrane complex (pm/oam), with some label attached particularly in regions with a “grazing” section plane, due to the permeabilization applied. This cell has been digitonized during aldehyde fixation for impregnation with primary AB and IgG-Au5 and embedded for incubation with the same ABs in sequence. Bar = 0.1 \mu m.
Assembly of F-actin around nascent phagosomes is well established, not only in mammalian cells but also in *Paramecium* cells (Allen and Fok 1983; Fok and Allen 1988). In detail, fusion of acidosomes with the nascent food vacuole depends on F-actin (Fok et al. 1987), as does maturation along the phago-lysosomal pathway, where multiple fusion/fission processes occur (Allen and Fok 1985; Allen et al. 1995). Interestingly, in our study, gold labeling immediately below the cytopharyngeal plasma membrane is less intense than between the closely packed globular and discoidal vesicles slightly below. This can be seen in line with the following reports. In *Dictyostelium*, F-actin prevents clustering of endosomal vacuoles (Drengk et al. 2003). Alternatively, in yeast, actin is required for Ca^{2+}-mediated vacuole interaction leading to fusion (Merz and Wickner 2004). The final step of this cycle in *Paramecium*, exocytotic release of spent phago-lysosomes, can also be inhibited by cytochalasin B (Allen and Fok 1985). In agreement with this previous work, the site of phagosome formation, vacuoles of different size, and the cytoproct are clearly labeled with anti-actin ABs in our CLSM and EM pictures. Therefore, the fine filaments described at the cytoproct by Cohen et al. (1984) are, at least to some extent, F-actin. However, centrin also occurs at the cytoproct, according to the CLSM pictures presented in Figure 3.

At the EM level, we see that the cytosolic compartment around large and small vacuoles is frequently heavily labeled (even when filaments are difficult to discern due to faint contrast resulting from preparation for immuno-EM analysis). This holds, e.g., for domains with clearly visible microtubule bundles deep inside the cell and for regions with discoidal vesicles approaching the cytopharynx. The latter are delivered along microtubule rails, using dynein as a motor (Schroeder et al. 1990). Therefore, actin at these sites may serve not as a motor, but rather as a kind of scaffold. In sum, apart from association with non-actin filaments (see below), we see that actin is also associated with the second cytoskeletal element, the microtubules. This agrees with functional data obtained by combined drug application (Fok et al. 1985).

Label also occurs around the oral cavity outside the site of phagosome formation in the cytopharynx. Such filaments are known not to represent actin, either in *Paramecium* (Clérot et al. 2001), or in other ciliates (Viguès et al. 1999). In these regions, F-actin may again serve structuring of these firmly established subcellular domains and/or vesicle trafficking. Interestingly, co-assembly of polymerizing actin with other filament components from *Tetrahymena* can be produced in vitro (Mitchell and Zimmerman 1985).

Vesicles deeper inside the cytoplasm, often close to a large phagosome, are also surrounded by gold label. All this reflects that actin is present throughout the cell in LM analyses, frequently as strands. Actin may thus participate directly or indirectly in vesicle trafficking, including cyclosis.

Not only ciliary basal bodies, but also the ciliary shaft, are labeled by anti-actin ABs. Labeling of cilia has been reported previously based on peroxidase-based preembedding immunostaining in *Paramecium* (Tiggemann and Plattner 1981) and in quail oviducts (Sandoz et al. 1982). Because this method is subject to redistribution artifacts (Plattner and Zingsheim 1983), we considered a re-analysis by Western blots and by the postembedding EM methodology to be necessary. It is known only from flagella of the green alga, *Chlamydomonas* (Mitchell 2000; Hayashi et al. 2001; Hirono et al. 2003), that actin is mandatory for normal beat activity. This may apply also to cilia of *Tetrahymena*, whose 14S axonemal dynein binds actin (Muto et al. 1994). More details on the role of actin in cilia remain to be elucidated.

Another poorly understood aspect concerns coupling of cortical calcium stores to the cell membrane. With mammalian cells, one of the molecules considered to establish such connections, particularly for store-operated Ca^{2+}-influx, is actin (Patterson et al. 1999; Rosado and Sage 2000; Kunzelmann-Marche et al. 2001; Wang et al. 2002). Interestingly, we find gold label that may be associated with the narrow subplasmalemmal space not only using a variation of the general labeling procedure that facilitates access of ABs (Figure 9), but also, though to a lesser extent, using postembedding labeling (Figures 5 and 7). This becomes evident particularly after statistical evaluation (Table 1). Although cytochalasin B application did not change concomittant Ca^{2+} signals (Mohamed et al. 2003), we keep this question open because the different actin isoforms found in *Paramecium* (Kissmehl et al., in preparation) may have different drug sensitivities.

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**Figure 10** Label around the oral cavity, in a region enriched in ciliary basal bodies (bb) in (A) or in vesicles (B). These represent, at least in part, discoidal vesicles (dv) known to recycle membranes for nascent phagosome formation, which may be assisted by round vesicles (rv) as discussed in the text. A particularly densely labeled domain in (A) is framed. Some label is located between unlabeled fibrous material (fm, not actin-type). Also note some label on basal bodies (bb) and within some cilia (ci). In (B), the label is scattered between the discoidal vesicles (dv). In (A) and (B), a &ge;1-μm-thick layer below the oral cavity plasma membrane is heavily labeled, starting at a distance from the plasma membrane. Bars = 0.1 μm.
Our present immunogold EM analysis largely depends on the preparation schedule used, whereas we obtained no such clear-cut labeling pattern with other approaches (data not shown). The current approach implied rapid injection (spraying) of cells in 0°C aldehyde fixative, containing high formaldehyde and very low glutaraldehyde concentrations, followed by low temperature embedding and UV polymerization at −35°C. This can considerably restrict diffusion of macromolecules and, even more, of filamentous aggregates. Therefore, we consider the current approach, elaborated on a (semi-)quantitative basis, more reliable than some previous attempts to localize actin in such cells.

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Literature Cited


Figure 11 Label of the cytosolic compartment around small vacuoles (sv), probably acidosomes, approaching a large one (lv), probably a nascent food vacuole (see text). Particularly labeled domains are highlighted (boxes 1, 2). Some of this label is associated with microtubule bundles (mt), e.g., in (B), but not in (A). In (B) a “trail” marked by box 2 is especially strongly labeled, although no distinct filament system can be recognized because of the preparation required. Only occasional gold grains are seen in non-cytosolic compartments, e.g., background in vacuole in (A). Bars = 0.1 μm.


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Actin in Paramecium