Actin Filament Organization in Activated Mast Cells Is Regulated by Heterotrimeric and Small GTP-binding Proteins

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Abstract. Rat peritoneal mast cells, both intact and permeabilized, have been used widely as model secretory cells. GTP-binding proteins and calcium play a major role in controlling their secretory response. Here we have examined changes in the organization of actin filaments in intact mast cells after activation by compound 48/80, and in permeabilized cells after direct activation of GTP-binding proteins by GTP-γ-S. In both cases, a centripetal redistribution of cellular F-actin was observed: the content of F-actin was reduced in the cortical region and increased in the cell interior. The overall F-actin content was increased.

Using permeabilized cells, we show that AlF₄⁻, an activator of heterotrimeric G proteins, induces the disassembly of F-actin at the cortex, while the appearance of actin filaments in the interior of the cell is dependent on two small GTPases, rho and rac. Rho was found to be responsible for de novo actin polymerization, presumably from a membrane-bound monomeric pool, while rac was required for an entrapment of the released cortical filaments. Thus, a heterotrimeric G-protein and the small GTPases, rho and rac, participate in affecting the changes in the actin cytoskeleton observed after activation of mast cells.

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permeabilized cells to GTP-γ-S. We find that a net centripetal redistribution of F-actin, observed after the activation of intact cells, could be mimicked simply by addition of GTP-γ-S to permeabilized cells in the absence of Ca2+ (pCa > 9). This effect results from a disassembly of the cortex and a de novo polymerization of actin in the cell interior. Using the permeabilized cell system to study these steps separately, we show that an AlF4–-sensitive heterotrimeric GTPase is involved in the disassembly of the cortical F-actin. In addition, we have examined the effects of two small GTPases, rho and rac, both of which have been implicated in the control of the cytoskeleton (see above), and show that both of them are required for the appearance of F-actin in the interior of the cell.

Materials and Methods

Materials

All nucleotides (GTP-γ-S, GTP, and ATP) were obtained from Boehringer Mannheim Biochemicals (Indianapolis, IN). SL-O was from Murex Diagnostics Ltd., Dartford, United Kingdom (cat. no. MR16, Lot K 908910). Vi4rhoA, Vi2ract, and Ni7ract proteins were purified as GST fusion proteins, subsequently cleaved and characterized as described (Ridley et al., 1992). Glass Multitest slides were from ICM-Flow. [32P]NAD was from NEN-DuPont (Boston, MA). C3 transferase purified from clostridium botulinum was a kind gift of Dr. Klaus Aktories (Rudolph-Buchheim-Institute fur Pharmacology, Giessen, Germany). All other reagents were obtained from the Sigma Chemical Co. (Poole, Dorset, United Kingdom).

Cell Preparation and Treatment

Preparation of rat peritoneal mast cells was as described previously (Cockcroft et al., 1987). The cells, in a solution containing 137 mM NaCl, 2.7 mM KCl, 1.0 mM CaCl2, 2 mM MgCl2, 56 mM glucose, 1 mg·mL−1 BSA, and 20 mM Na-Pipes, pH 6.8 (chloride buffer [CB]), were allowed to attach to glass Multitest slides (6-mm diameter wells) for 1 h at room temperature. Intact cells were stimulated with 5 μM of compound 48/80 in CB for 2 min at 30°C and were then fixed for 20 min with 3.8% formaldehyde in phosphate-buffered saline containing 2 mM MgCl2 and 3 mM Na-EGTA (PBSME). To permeabilize the attached cells, the CB was exchanged for 137 mM Na-glutamate, 2 mM MgCl2, 1 mg·mL−1 BSA, 20 mM Na-Pipes, pH 6.8 (glutamate buffer [GB]), and the cells were then exposed for 8 min to streptolysin-O at 0.4 IU·mL−1 in GB containing 3 mM Na-EGTA (GB-EGTA) at room temperature. Permeabilized cells were then washed free of soluble components and excess SL-O with GB-EGTA. 10 min after SL-O addition, cells were treated for 15 min with 30 μM GTP-γ-S (or AlF4–, or 2 μg·mL−1 Vi4rhoA or 2 μg·mL−1 Vi2ract) in GB-EGTA at 30°C and then fixed. The 15-min incubation is sufficient to ensure the completion of the secretory response (Koffer, 1993).

AlF4– was obtained by addition of 30 mM NaF and 10 μM AICl3 to glutamate buffer in the absence of EGTA (chelation of the aluminum ions with EGTA or EDTA rendered the fluoride treatment ineffective). The unbuffered calcium levels (~1 μM), resulting from the omission of EGTA at this stage, did not affect the cytoskeleton, provided that the cells were initially permeabilized in the presence of the chelator. To obtain recombinant Vi4rhoA or Vi2ract in the GTP form, the proteins (73 μg·mL−1) were incubated with 100 μM GTP in 50 mM Tris containing 5 mM EDTA, pH 7.0, for 20 min, followed by the addition of 10 mM MgCl2. Controls were exposed to the appropriate final concentrations of GTP, Tris, or EDTA (2.7 μM, 1.4, and 0.1 mM, respectively). Vi4rhoA was active only in the GTP form, prebinding of the protein with GDP rendered it inactive. 0.1 μg·mL−1 C3 transferase was added at the time of permeabilization in the presence of 0.5 mM NAD+, maintained throughout the wash, and removed just before the addition of GTP-γ-S (i.e., 10-min treatment). Nitracell was also included with the SL-O and wash at a concentration of 8 μg·mL−1, but was additionally maintained throughout the incubation with GTP-γ-S. 10 μM cytochalasin E was added 3 min after permeabilization and was maintained throughout treatment with GTP-γ-S or Vi4rhoA.

Freely soluble ATP would be expected to leak from SL-O-permeabilized cells. However, to ensure ATP depletion, in some experiments, the cells were metabolically inhibited. This was achieved by incubation of the attached cells with 6 mM 2-deoxyglucose and 10 μM antimycin A in GB at 37°C for 7 min before the addition of SL-O.

F-Actin Staining Protocols

Postlabeling. Total cellular F-actin was visualized by staining-fixed cells for 15 min with 0.6 μM rhodamine phalloidin (RP) in PBSME containing 80 μg·mL−1 lysophosphatidyl choline (L-4129; Sigma Chemical Co.).

Prelabeling. To follow the movement of existing F-actin filaments, the cells were labeled before the addition of GTP-γ-S. 4 min after permeabilization, either 0.18 or 2.0 μM RP in GB-EGTA was added. The stain was thoroughly washed off 2 min before the addition of GTP-γ-S (i.e., 4-min exposure to RP). After the usual 15-min exposure to GTP-γ-S, the cells were fixed as described above.

Pre (2.0 μM RP) - and Postlabeling. This protocol was performed to stabilize the F-actin cortex before the addition of GTP-γ-S and then to visualize any additional F-actin formed after the treatment with the nucleotide. The cells were prelabelled with 2.0 μM RP, and the stain washed off as for prelabeling. GTP-γ-S was added for 15 min, and the cells fixed and restained with RP as for postlabeling.

Microscopy and Flow Cytometry

Fixed and stained cells were mounted in PBSME (so glycerol was used because this can lead to cell collapse) and fluorescent images were obtained using a confocal laser scanning microscope (Leica Inc., Deerfield, IL) attached to a microscope (Fluovert-FU; E. Leitz, Inc., Rockleigh, NJ). Excitation was at 514 nm, and emission was >595 nm. Digital images (equatorial optical slices) are displayed using the Leica "grau" look-up table. The equatorial plane of focus was defined as the midpoint between the top and the bottom of the cell. The nucleus usually, but not always, lies on this plane.

Image Analysis

F-Actin Content. F-actin content was determined as a mean of the total pixel RP intensity per equatorial slice (n > 50) and related to the appropriate control level. This was confirmed by an "extended focus" method: an image stack was obtained, encompassing the entire depth of the cell, by superposition of ~20 optical slices, each 1 μm apart, along the cell's z axis. The relative increase in F-actin content after 48/80 or GTP-γ-S treatment obtained by this method was not significantly different from that obtained by the analysis of the area of equatorial slices. The equatorial plane of focus was defined as the midpoint between the top and the bottom of the cell.

F-Actin Distribution. Images of equatorial slices were quantified by radial line scan analysis (as shown in Fig. 4). Three lines were obtained per cell for a total of ≥50 cells and the RP fluorescence intensity profiles for each set of experimental conditions were pooled and averaged to form a mean profile. Data obtained from these scans were further analyzed to obtain bar charts (see Figs. 5 and 7). The cortical and the interior regions were designated as regions encompassed by pixel numbers 1-10 and 11-30, respectively. These represent distances of 0-2.45 and 2.45-7.35 μm from the cell edge. Changes in the RP staining of these two regions were calculated as percentages of the total RP intensity in both regions from appropriate control cells (i.e., the sum of intensities of pixels 1-30). The nucleus was normally >8 μm from the cell edge and, therefore, not included in this analysis.

Leakage of F-Actin

Mast cells were attached to 13-mm diameter glass coverslips (at a density of 2×106 cells·mm−2) placed within 15-mm wells and permeabilized as described above. At various times after the addition of SL-O, the permeabilizing solution was aspirated from the attached cells. The supernatant and cells were added to sample buffer (Laemmli, 1970) and boiled for 5 min. Determination of actin in these extracts was by SDS-PAGE, followed by densitometry of silver-stained gels as described previously (Koffer and Comports, 1989).

[32P]ADP Ribosylation with C3 Transferase

Cells were attached and permeabilized as described above for leakage of ac-
tin. In addition, 4.5 μCi [32P]NAD/well (0.5 μM) and 0.1 μg·ml⁻¹ C3 transferase were included in the permeabilization buffer. At various times, this supernatant (containing the bulk of the radioactivity) was discarded, and the remaining cells were extracted with the sample buffer and analyzed by a 15% SDS-polyacrylamide gel (Laemmli, 1970). The gel was dried, and labeled proteins were visualized using phosphor-imaging plates and Bio-Imaging Analyzer FUJIX BAS1000 (Fuji, Tokyo, Japan).

The proportion of soluble rho was determined in suspended cells. Mast cells in CB were preincubated on ice with SL-O (0.8 IU·ml⁻¹) for 5 min (no permeabilization occurs at this temperature), centrifuged at 4°C, and then resuspended in aliquots (250,000 cells in 200 μl) in GB containing 1 mM phenyl-methylsulphonyl fluoride, 10 μg·ml⁻¹ leupeptin, 10 μg·ml⁻¹ pepstatin, 1 mM EDTA, 1 mM EGTA, and 0.5 mg BSA·ml⁻¹. Permeabilization was initiated by incubating the cells at 30°C, and samples were withdrawn after 10 and 30 min and centrifuged (1 min at 12,000 g). The pellets were resuspended in 200 μl of the above buffer, and both supernatants and pellets were incubated for 20 min at room temperature with C3 transferase (0.1 μg·ml⁻¹) and 3.7 μCi [32P]NAD (2.5 μM). Proteins were then precipitated by addition of 25 μl 100% trichloroacetic acid (30 min on ice), pelleted (2 min at 1,200 g), 2× washed with ethanol-ether (1:1), dissolved in 50 μl of sample buffer, and analyzed as above.

Results

Response of Intact Cells to Compound 48/80

RP staining of quiescent mast cells revealed F-actin to be present primarily at the cortex (Fig. 1A). After stimulation of intact mast cells by compound 48/80, this pattern changed (Fig. 1B); the staining of the cell cortex was now less intense, while that of the cell interior became stronger. There was a marked (~30%) overall increase in the total F-actin

![Figure 1](image-url)

**Figure 1.** Confocal micrographs of rhodamine phalloidin–stained mast cells showing F-actin distribution in (A) unstimulated and (B) 48/80-treated intact mast cells. (C) Unstimulated and (D) GTP-γ-S–treated, permeabilized cells showed similar patterns. Different patterns were exhibited by permeabilized cells treated with (E) AlF₄⁻ and (F) V14rhoA. (G) shows the internal structures obtained after treatment with GTP-γ-S at higher magnification. The bars in A and G represent 20 and 10 μm, respectively.
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by confocal microscopy and flow cytometry, respectively.

48/80. In spite of the extensive leakage of the monomer,
was filled with polymerized actin. Identical changes
futures that appear as if the intergranular cytoplasmic space
were also obtained in experiments where the interval be-
content (Fig. 2), as measured by confocal microscopy or

Flow cytometry.

Leakage of Actin from Streptolysin-permeabilized Cells

In the experiments described below, cells attached to glass
slides were permeabilized by addition of SL-O (0.4 IU.mL^-1)
and washed to remove any endogenous ions, nucleotides,
and freely soluble components (Koffer, 1993). To establish
the proportion of actin remaining within these cells, we
determined the extent and time course of actin leakage. Fig. 3
shows that 70-80% of total cellular actin leaked out after
permeabilization, and that this process was complete within
4-8 min. Subsequent washing of permeabilized cells did not
remove any further actin (not shown). The strong cortical
staining of F-actin was not reduced after the permeabiliza-
tion (compare Fig. 1 a and c), indicating that the leakage is
caused by the diffusion of actin monomers rather than loss
of filaments after cytosol dilution. This is consistent with
previous results obtained with mast cells in suspension
(Koffer et al., 1990) and with other cell types (Cassimeris
et al., 1990), which show that the cortical F-actin is very
resistant to depolymerization despite the removal of the solu-
ble G-actin pool.

Response of Permeabilized Cells to GTP-γ-S

GTP-γ-S was introduced into washed permeabilized cells 10
min after the addition of SL-O. When only a negligible frac-
tion of the (freely diffusible) G-actin pool remains within the
cells. The Ca^2+ concentration was clamped below 1 nM by
EGTA. As shown in Figs. 1 D and 2, addition of this nucleo-
tide was sufficient to produce changes in the actin cytoskele-
ton indistinguishable from those induced in intact cells by
48/80. In spite of the extensive leakage of the monomer,
F-actin content increased by ~23% and 19% as measured
by confocal microscopy and flow cytometry, respectively
(Fig. 2). Higher magnification of cells stimulated by GTP-
γ-S (Fig. 1 G) or 48/80 (not shown) showed internal struc-
tures that appear as if the intergranular cytoplasmic space
has been filled with polymerized actin. Identical changes
were also obtained in experiments where the interval be-
tween the permeabilization and the addition of GTP-γ-S was
longer (<20 min).

No ATP was required for this response. To confirm this,
we have depleted the endogenous ATP content by a pretreat-
ment of cells with metabolic inhibitors, deoxyglucose and
antimycin A, before permeabilization. Exposure of these
cells to GTP-γ-S produced similar results (data not shown).
Furthermore, to exclude the participation of protein kinase
C, we have also treated the cells with a protein kinase C in-
hibitor, H7 (100 μM, added at the time of permeabilization
and maintained throughout GTP-γ-S treatment) (Kawamoto
and Hidaka, 1984). Again, there was no significant effect on
the response.

Quantitative Analysis of Changes in F-Actin Distribution

Changes in the distribution of F-actin after the stimulation of
intact and permeabilized cells were further analyzed. Im-
ages, such as those shown in Fig. 1, were quantified by radial
line scan analysis of equatorial slices, and the RP fluores-
cence intensity profiles for each set of experimental condi-
tions were pooled. The cortical and the internal regions were
designated as regions encompassed by pixel numbers 1-10 and
11-30, respectively. These represent distances of 0-2.45
and 2.45-7.35 μm from the cell edge (Fig. 4 B). The nucleus
was normally >8 μm from the cell edge and, therefore, not
included in this analysis. Fig. 4 A confirms and quantifies the
effects of GTP-γ-S seen in Fig. 1 D: a selective decrease and
increase of F-actin presence in the cortical and the internal
regions, respectively. Very similar scans were obtained for
stimulated intact cells.

Data obtained from these scans were expressed in the form
of a bar chart shown in Fig. 5. Changes in the regions desig-
nated as cortical (full bar) and internal (hollow bar) were cal-
culated as percentages of the total F-actin content in both

Figure 2. Relative F-actin content of intact and permeabilized mast
cells after treatment with compound 48/80 and GTP-γ-S. Values
were obtained by quantification of individual cells from confocal images similar to those in Fig. 1 and expressed as percentage of
control. The bars with asterisks (*) show confirmation of these
values by flow cytometry. Values obtained from confocal imaging
are mean ± SEM, n > 50 cells. Each asterisked bar represents the
mean of two experiments, each involving ≥5,000 cells.

Figure 3. Time course of actin leakage from SL-O-permeabilized
mast cells. Cells were attached onto glass coverslips. The actin re-
maining within the attached cells and that leaking into the superna-
tant was determined at various times after the addition of strep-
tolysin-O (0.4 IU.mL^-1) and plotted as a percentage of total
cellular actin content. Determination of actin was by SDS-PAGE,
followed by densitometry of silver-stained gels. The time at which
GTP-γ-S was routinely added to the incubation (to perform experi-
ments presented in Figs. 1, 3, 4, and 5) is shown by the arrow.
regions from appropriate control cells (the sum of intensities of pixels 1–30). Thus, relative changes in distribution of F-actin induced by GTP-γ-S could be easily evaluated (Fig. 5, group II, first pair): a reduction in the cortical region of ~10% and an increase in the interior region of ~30%. Similar changes were produced by compound 48/80 in intact cells, except that the increase in the cell interior region was slightly greater (Fig. 5, group I).

The Fate of the Cortical Filaments
To examine the fate of existing filaments, the cells were
The distribution change shown in Fig. 4 a seems to result from disassembly of cortical F-actin combined with movement of filaments from the cortex to the cell interior, together with a de novo polymerization of actin from a G-actin pool that is apparently retained within permeabilized cells. However, it is also possible that previously "hidden" filaments (such as would have been inaccessible to RP) become accessible to RP after GTP-γ-S treatment. We have tested the effects of cytochalasin E, a microfilament-destabilizing agent that blocks filament barbed ends (Cooper, 1987). This agent prevented the GTP-γ-S induced overall increase in F-actin content (not shown), indicating de novo actin polymerization from newly exposed barbed ends rather than exposure of hidden RP-binding sites.

**Activation of Heterotrimeric G Proteins by AIF:**

Causes Cortical Disassembly

GTP-γ-S affects the activity of both heterotrimeric and small, ras-related, GTP-binding proteins. To differentiate between the contributions of these two classes, we first tested the effects of AIF, a specific activator of trimeric G-proteins (Kahn, 1991). Treatment of permeabilized cells with AIF led to selective loss of cortical F-actin (Figs. 1 e; 5, group III). This indicates that a heterotrimeric G-protein is directly involved in disassembly of the actin cortex. However, no increase in F-actin in the cell interior was observed. Thus, activation of this AIF-sensitive G-protein was not sufficient to relocate cortical filaments to the cell interior neither could it induce the de novo actin polymerization. Activity of additional GTP-binding proteins must, therefore, be required for the entrapment of the released filaments and for the de novo actin polymerization.

**A Substrate for the C3 Transferase is Present in Permeabilized Mast Cells**

Members of the rho family of GTP-binding proteins are the most obvious candidates for the above roles. These proteins are substrates for the C3 transferase from clostridium botulinum. This enzyme inactivates rho proteins by ADP ribosylation (Aktories et al., 1989). To test for the presence of rho family proteins, mast cells were permeabilized with SL-O in the presence of C3 transferase and [32p]NAD for varying lengths of time. The radioactive proteins remaining in the cells were then visualized by SDS-PAGE followed by phosphor imaging. The results (Fig. 6) show one major radioactive band at 20 kD, the ribosylation of which reached a maximum after 10 min of incubation with C3. No ribosylation was evident in the absence of the toxin.

To establish the proportion of soluble and membrane-bound rho, leakage experiments were performed on cells in suspension. Results are shown in Table I. Only ~40% of C3 substrate (the 20-kD band) was found to leak out of the cells after their permeabilization, and the leakage was complete within 10 min. Using a recombinant rho protein as a standard, ~0.85 ng ± 0.4 (n = 3) of rho (total) was calculated to be present per 100,000 cells. This represents ~0.3 μM rho, when extrapolated to intact cells (cell volume is ~1.3 x 10^-12 liters).

**Rho Induces Polymerization of Actin in the Cell Interior**

Having established the presence of rho proteins remaining within mast cells after permeabilization, the constitutively active recombinant V14rhoA mutant (Ridley and Hall, 1992; Ridley et al., 1992) was used to investigate any involvement...
Ribosylation of rho by C3 transferase. Attached mast cells were permeabilized with SL-O in the presence of C3 transferase and [3P]NAD. After the indicated times, the cell supernatant was removed, and radioactive proteins remaining within the cells were analyzed by SDS-PAGE and phosphor-imaging. Lanes 2-5 show treatment with C3 for 20, 15, 10, and 5 min, respectively. A protein of ~20 kD is ribosylated in a time-dependent fashion, reaching a maximum after 10-15 min. Lane 1 shows that no radioactive bands are obtained in the absence of C3.

Table I. Leakage of C3 Transferase Substrate from Permeabilized Mast Cells

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>PSL</th>
<th>Percentage (P+S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 min</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pellet (P)</td>
<td>4,893</td>
<td>56</td>
</tr>
<tr>
<td>Supernatant (S)</td>
<td>3,901</td>
<td>44</td>
</tr>
<tr>
<td>P+S</td>
<td>8,794</td>
<td></td>
</tr>
<tr>
<td>30 min</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pellet (P)</td>
<td>5,985</td>
<td>62</td>
</tr>
<tr>
<td>Supernatant</td>
<td>3,691</td>
<td>38</td>
</tr>
<tr>
<td>P+S</td>
<td>9,676</td>
<td></td>
</tr>
<tr>
<td>Unfractionated</td>
<td>8,317</td>
<td></td>
</tr>
</tbody>
</table>

ADP-ribosylation with [3P]NAD was performed as described in Materials and Methods. Proteins were separated by SDS-PAGE (100,000 cells/lane) and analyzed by a Bio-Imaging Analyzer. Results are expressed as phosphosensitive luminescence intensity obtained from appropriate bands (PSL, background subtracted).

Rho is not Required for Relocalization of Cortical Filaments

Fig. 7 shows changes induced by the addition of GTP-γ-S in the presence of various agents. These changes are shown in two ways: (a) the overall F-actin distribution change ("post"), which is the sum of the filament disassembly, relocalization, and the de novo actin polymerization (as determined by post-labeling experiments described in Fig. 4 a); and (b) the change caused by the filament relocalization only ("pre," as determined by pre-labeling experiments described in Fig. 4 b). Inhibition of rho by C3, or blocking of barbed ends by cytochalasin E, produced identical effects. In the presence of either of these agents, GTP-γ-S failed to induce an increase in cellular F-actin content. However, the relocalization of cortical filaments to the cell interior was unaffected, showing that the nucleation sites activated by rho are not responsible for entrapment of filaments, liberated from the cortical region by the AlF₄⁻-sensitive G-protein. In some experiments, C3 and cytochalasin caused a fall in the resting levels of F-actin. The reduction, however, was never >20%, and...
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it became less apparent after increasing the time between permeabilization and the addition of these agents.

**Inhibition of the Activity of rac by N17racl Prevents Filament Relocalization**

Since neither heterotrimeric G-proteins nor rho were involved in filament relocalization, we tested the role of rac, a rho-related protein. N17racl has been used previously in microinjection studies as a dominant negative inhibitor of rac (Ridley et al., 1992). Fig. 7 shows that addition of this protein to permeabilized cells completely blocks GTP-γ-S-induced filament redistribution from the cortex to the cell interior. Using the prelabeling protocol, the cortical staining was reduced by addition of GTP-γ-S, but there was no commensurate increase in cell interior staining. This indicates that rac is required for the entrapment of the released cortical filaments within the cell interior. De novo actin polymerization, however, was unaffected by N17racl because a considerable increase in internal staining was still detected by the postlabeling protocol (Fig. 7). In addition, N17racl did not affect the basal levels of cellular F-actin (not shown). No changes in the cytoskeleton were observed upon addition of the constitutively active form of rac, V12racl (not shown). It appears that in the absence of cortical actin disassembly, no relocation of filaments can occur.

**Inhibition of both rho and rac Causes GTP-γ-S-dependent Loss of Filaments**

Since the activities of rho and rac are required for actin polymerization and filament entrapment, respectively, it follows that inhibition of both processes together should cause a reduction in total cellular F-actin content in response to GTP-γ-S. Fig. 7 shows that this is in fact the case. In the presence of N17rac together with either C3 or cytochalasin E, addition of GTP-γ-S leads to a significant loss of F-actin from the cell (both of pre- and postlabeled filaments). This is similar to the situation observed after the addition of AlF₄⁻ to permeabilized cells (Fig. 5, group III). In both cases, presumably, there is no filament entrapment or actin polymerization because rho and rac are not activated.

**Discussion**

**GTP-binding Proteins Regulate Mast Cell Microfilaments Independently of Freely Soluble Cytosolic Components**

We show here that GTP-binding proteins play a major role in the cytoskeletal reorganization that accompanies mast cell activation. Two changes in F-actin morphology were elicited by the exposure of permeabilized cells to GTP-γ-S: the disassembly of the cortex and the appearance of actin filaments in the cell interior. Released cortical filaments, as well as newly polymerized actin, were recruited into this interior cytoskeleton. These effects result (at least in part) from the GTP-γ-S activation of two small GTPases, rho and rac, and of a heterotrimeric G-protein(s).

GTP-γ-S-induced reorganization of actin proceeded in the absence of any freely soluble cytosolic components, indicating that the GTP-binding proteins and any accessory proteins must be associated with intracellular structures and remain within mast cells for >10 min after SL-O permeabilization. This is supported by our finding that the inhibitors of both rho and rac had profound effects on the response of the permeabilized cells. These results are somewhat unexpected because in the presence of Mg²⁺, the binding of GTP-γ-S to small GTPases requires a nucleotide exchange protein (Downward, 1990; Hall, 1993; Takai et al., 1992). Exchange proteins specific for rho and rac have not been identified so far (Hiraoka et al., 1992). The exchange factor for a small GTPase ras, sos, has been found to be translocated from the cytosol to the plasma membrane upon activation of a tyrosine kinase receptor, resulting in a formation of a multiprotein complex that includes the activated ras (Egan et al., 1993). Sos, however, does not leak from quiescent cells when they are permeabilized with SL-O (Downward, J., personal communication). Using cell fractionation studies, rac has been found to be cytosolic in quiescent neutrophils, and to translocate to the plasma membrane upon exposure of the cells to activating stimuli, again participating in the formation of a complex of the NADPH-oxidase system (Abo et al., 1991; Quinn et al., 1993). Similarly, thrombin activation of platelets resulted in an increased association of rho with cytoskeletal fractions (Zhang et al., 1993). We have found the major part (~60%) of rho (as a C3 substrate) to be associated with internal structures of (resting) mast cells (Table I). It is possible that small GTPases and their exchange factors are loosely associated with the appropriate membrane, adjacent to their effectors. Signals, which activate the cell, would then merely consolidate this association and promote the assembly of a multiprotein complex.

Since freely soluble endogenous nucleotides have also been removed during the permeabilization and washing procedures, and no exogenous ATP is required, it is unlikely that a phosphorylation event constitutes an integral part of the GTP-γ-S-induced response. Moreover, neither depletion of ATP before the addition of SL-O (by incubating cells with metabolic inhibitors) nor treatment of permeabilized cells with H7, an inhibitor of protein kinase C (Kawamoto and Hidaka, 1984), affected the response. However, addition of MgATP enhanced the proportion of permeabilized cells that responded to GTP-γ-S, and it caused an increased F-actin presence in the cell interior (data not shown).

It is interesting to note that the centripetal rearrangements of the mast cell cytoskeleton occurred in the absence of Ca²⁺. Previous results have shown that Ca²⁺ introduced into permeabilized mast cells, can cause a substantial loss of F-actin content independently of GTP-γ-S (Koffer et al., 1990). Thus, it appears that two routes to cytoskeletal regulation, one independent of and the other dependent on Ca²⁺, operate in mast cells and most probably interact and modify each other. The absence of Ca²⁺, ATP, and the cytosolic factors would exclude the involvement of most second messengers, but a G-protein–induced change in phospholipid composition may well mediate some of these effects. In this context, recent evidence for the activation of both phosphatidylinositol 3-kinase (Zhang et al., 1993) and phospholipase D (Bowman et al., 1993) by rho related protein may be relevant.

**Disassembly of Cortical Microfilaments**

AlF₄⁻ treatment of permeabilized mast cells could mimic one aspect of the GTP-γ-S–induced response. It resulted in a selective decrease of cortical F-actin, indicating a presence.
of an heterotrimeric G-protein that regulates disassembly of cortical actin filaments and/or their detachment from the plasma membrane. Higher (> μM) concentrations of RP protected the cortical filaments from both GTP-γ-S or AIF₄⁻-induced disassembly. Thus, a calcium independent, phalloidin-sensitive and membrane-associated, actin-severing protein may be involved. A possible candidate is actin depolymerizing factor, whose severing activity is inhibited by phalloidin (Maciver et al., 1991) and regulated by phosphoinositides (Yonezawa et al., 1990), and whose association with plasma membrane has previously been demonstrated (Koffer et al., 1988).

The response of neutrophils to both GTP-γ-S and AIF₄⁻ has been studied previously and found to be somewhat different from that of mast cells. In electropereamabilized neutrophils, AIF₄⁻ and GTP-γ-S both caused an increase in actin content (Therrien and Naccache, 1989; Bengtsson et al., 1990). This may be caused by the much smaller lesions in electropereamabilized cells, which result in the retention of the bulk of the monomeric G-actin pool and cytosolic components.

**De Novo Actin Polymerization in the Cell Interior**

The appearance of new filaments in the cell interior was partially reproduced by addition of V14rhoA to permeabilized cells. Newly polymerized filaments appeared despite the loss of soluble actin. New filament formation was sensitive to cytochalasin E and, therefore, dependent on the presence of free filament barbed ends. The implications of this are important, suggesting the existence of a membrane-bound (or slow-leaking) G-actin pool that can be readily mobilized when actin polymerization is required. Thus, rho may initiate the release of this sequestered G-actin together with the formation of new actin nucleation sites.

Recent evidence suggests that such membrane-associated G-actin pool may well exist (Herman, 1993; Cao et al., 1993). This sequestered pool of G-actin could represent a ready-to-go pool of monomer stored where it is most needed, near the advancing edge of a migrating cell. The increase in F-actin content of mast cells was quantitatively similar, regardless of whether the cells were intact or permeabilized. This suggests that the same membrane-bound pool of G-actin may also be recruited after activation of intact cells by 48/80.

Examination of the high magnification micrograph (Fig. 1 G) suggests that actin polymerization after exposure to GTP-γ-S (or V14rhoA) occurs on the secretory granule membranes which may, therefore, be the location of rho in these cells. Indeed, C3 substrates on purified neutrophil granule membranes have already been identified (Philips et al., 1991). The role of rho in stress fiber production has been postulated to be via stabilization of a multiprotein complex, components of which are recruited from the cytosol, creating focal adhesion from which the fibers emanate (Ridley and Hall, 1992). In mast cells, the F-actin, which was seen to form around secretory granules, may also originate from multiprotein complexes that may provide new (cytochalasin sensitive) nucleation sites. However, in mast cells, the components of such a complex must already be associated with the appropriate (internal) membranes because they are resistant to leakage after permeabilization.

**Relocalization of Cortical Filaments**

In the presence of GTP-γ-S, disassembled cortical filaments did not leave permeabilized cells, and they were redistributed to their interior (Fig. 5, group II). On the other hand, after the AIF₄⁻ treatment, the filaments were lost from cells (Figs. 1 E and 5, group III), presumably because of their depolymerization into monomers and/or short oligomers and leakage. The experiments using N17rac imply that activation of rho is required to entrap the filaments within the cell interior. This entrapment was unaffected by C3 exoenzyme or cytochalasin E, and thus does not involve rho nor a barbed end-on polymerization event. The mechanism of this presumably rho-dependent entrapment is unknown; a side-on or a pointed end-on (cytochalasin insensitive) association of the filaments with internal membranes and/or actin may be responsible. Rac may also act to stabilize the released cortical filaments, preventing their depolymerization. A number of actin-associated proteins are known to bind to the sides of F-actin, for instance tropomyosin, α-actinin and actin-binding protein, which can act to stabilize the filaments or promote parallel and orthogonally cross-linked arrays of actin, respectively (Pollard and Cooper, 1986). Rac activity has been shown to be necessary for the formation of membrane ruffles in fibroblasts (Ridley et al., 1992). Membrane ruffles are rich in F-actin, which is extensively cross-linked. The role of rac in fibroblasts may be not only to initiate actin polymerization, but also to stabilize and cross-link filaments within ruffles while reducing the rigidity of the cortex.

No effect was observed following the introduction of an activated rac (V12rac) into permeabilized cells. This is most probably caused by the absence of cortical actin disassembly and, therefore, the absence of any filaments to be relocalized. In addition, no freely soluble pool of actin monomer is present in permeabilized cells, such as may have been required for the formation of ruffles in fibroblasts.

**Conclusions**

Treatment of intact cells with compound 48/80 produced changes in microfilament morphology, which were very similar to those induced by GTP-γ-S in permeabilized cells: a centripetal rearrangement of F-actin. The similarity of these responses supports the evidence for a direct (i.e., receptor-independent) activation of heterotrimeric G-proteins by compound 48/80 and other basic secretagogues (Mousli et al., 1990; Aridor et al., 1990, 1993). To transduce signals from this 48/80-sensitive, plasma membrane-associated, heterotrimeric G-protein to rho and rac in the cell interior, we postulate the existence of a cytosolic factor(s). This is illustrated in the scheme shown in Fig. 8. These cytosolic components would be lost from permeabilized cells, and this may be the reason why the AIF₄⁻ effect is limited only to the cortical disassembly.

We conclude that a heterotrimeric G-protein releases filaments from the cortex, which are then trapped within the cell by a rho-dependent mechanism. In addition, rho mobilizes a membrane-associated G-actin pool and mediates polymerization of actin in the cell interior. Together, these three activated GTP-binding proteins can induce centripetal rearrangements of the mast cell cytoskeleton independently of Ca²⁺.
Centripetal movement of actin filaments is a phenomenon that has been reported in many cell types and accompanies cellular processes such as locomotion or pseudopodial extensions (Heath, 1983; Theriot and Mitchison, 1992). This study provides a quantitative insight into another example of such movements in model secretory cells. The relevance of these cytoskeletal changes to the exocytotic process remains to be established. Since GTP-γ-S-induced exocytosis is fully inhibited by 1 mM GDP-β-S (Koffer, 1993), whereas inhibition of cytoskeletal changes requires higher (10 mM) concentrations (not shown), the G-protein(s) postulated to control exocytosis in mast cells, Ga (Cockcroft et al., 1987), cannot be responsible for this type of cytoskeletal redistribution. This study is, however, consistent with the postulate that cortical disassembly is a necessary preliminary step for secretion (Burgoyne and Cheek, 1985; Linstedt and Kelly, 1987). Possible roles for internal filaments, on the other hand, could be to provide a structural support for degranulating cells and to restrict granule movement, thus tempering any uncontrolled secretory response.

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