Protein phosphatases 1 and 2A transiently associate with myosin during the peak rate of secretion from mast cells.

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ABSTRACT

Mast cells undergo cytoskeletal restructuring to allow secretory granules passage through the cortical actomyosin barrier to fuse with the plasma membrane and release inflammatory mediators. Protein phosphorylation is believed to regulate these rearrangements. Although some of the protein kinases implicated in this phosphorylation are known, the relevant protein phosphatases are not. At the peak rate of antigen-induced granule mediator release (2.5 min), protein phosphatases PP1 and PP2A, along with actin and myosin II are transiently relocated to ruffles on the apical surface and a band at the peripheral edge of the cell. This leaves an area between the nucleus and the peripheral edge significantly depleted (3-5 fold) in these proteins. PMA plus A23187 induce the same changes, at a time coincident with its slower rate of secretion. Co-immunoprecipitation experiments demonstrated a significantly increased association of myosin with PP1 and PP2A at the time of peak mediator release, with levels of association decreasing by 5 min. Jasplakinolide, an inhibitor of actin assembly, inhibits secretion and the cytoskeletal rearrangements. Surprisingly, jasplakinolide also affects myosin, inducing the formation of short rods throughout the cytoplasm. Inhibition of PP2A inhibited secretion, the cytoskeletal rearrangements and led to increased phosphorylation of the myosin heavy and light chains at PKC-specific sites. These findings indicate that a dynamic actomyosin cytoskeleton, partially regulated by both PP1 and PP2A is required for mast cell secretion.
INTRODUCTION

The cross-linking of receptor bound IgE on the mast cell surface, triggers a sequence of intracellular events that culminate in the extracellular release of potent inflammatory mediators, many of which are stored in the secretory granules (Beaven and Baumgartner, 1996; Razin et al., 1995). Conversely, mast cell granule mediators, especially TNF-α have also been shown to play a central role in the defence against bacterial infection (Echtenacher et al., 1996; Malaviya et al., 1996). Therefore, understanding the mechanism through which mast cells release these mediators is of broad physiological importance.

In mast cells, the proteins regulating the heterotypic fusion process between the secretory granules and plasma membrane have been identified as SNARE related proteins (Guo et al., 1998; Hibi et al., 2000). However, the granules in mast cells are kept apart from their fusion sites on the plasma membrane by a cortical cytoskeletal barrier made up predominantly of actin and myosin. How the granules are moved towards and through this actomyosin barrier to their fusion sites is not understood in any regulated exocytotic cell. Recent studies in a variety of secretory cells, using inhibitors of actin or myosin function, have shown that remodelling of the actomyosin cortex is a prerequisite for regulated exocytosis (Lang et al., 2000; Valentijn et al., 2000).

Exocytosis is accompanied by distinct morphological changes that are believed to facilitate the removal or rearrangement of the cortical barrier (Lang et al., 2000; Oheim and Stuhmer, 2000; Sullivan et al., 2000; Valentijn et al., 2000). In activated mast cells these include the formation of ruffles or lamellae on the apical surface that contain both actin and myosin (Ludowyke et al., 1994a; Pfeiffer et al., 1985; Sahara et al., 1990) and the spreading of the cells upon the underlying substratum (Ludowyke et al., 1994a; Oliver et al., 1988). Correlated with these changes are increased assembly of F-actin (Apgar, 1991; Oliver et al., 1988) the formation of focal adhesions (Kawasugi et al., 1995) and actin and myosin-
containing plaques and filaments on the basal surface (Ludowyke et al., 1994b; Pfeiffer and Oliver, 1994).

The actomyosin cortical web is dynamic, being rapidly disassembled and assembled in response to extracellular signals, being controlled in part by the phosphorylation of myosin at specific sites on the heavy and light chains (Bresnick, 1999; Sellers, 1991). In contrast to smooth muscle, myosin IIA is the only conventional myosin isoform found in RBL-2H3 cells (Choi et al., 1996). In unstimulated cells, the myosin light chain (MLC) is phosphorylated by myosin light chain kinase (MLCK) and the myosin heavy chain (MHC) is phosphorylated at a number of sites by unknown kinases (Ludowyke et al., 1989). Activation of these cells with antigen or calcium ionophore does not affect phosphorylation at these sites, but leads to the phosphorylation of both the heavy chains and light chains by protein kinase C (PKC) at distinct sites (Choi et al., 1994; Ludowyke et al., 1989; Ludowyke et al., 1996). Recent work has shown that the MHC is also phosphorylated by CaM kinase II during activation (Buxton and Adelstein, 2000).

The balance of specific kinases and phosphatases dynamically regulates protein phosphorylation. The serine and threonine phosphatases are separated into 2 main families, defined as PPP and PPM. The predominant members of the PPP family are protein phosphatase type 1 (PP1), PP2A and PP2B (calcineurin) while PP2C is the predominant PPM protein phosphatase. PP1 and PP2A are considered the major intracellular phosphatases, which together account for >90% of all serine/threonine dephosphorylation reactions (Cohen, 1989; Cohen and Cohen, 1989; Mumby and Walter, 1993; Wera and Hemmings, 1995). Whilst the phosphatases which regulate myosin in mast cells are not known, in muscle cells it is clear that the primary phosphatase effecting the MLC is PP1 (Chisholm and Cohen, 1988; Hartshorne et al., 1998). This phosphatase dephosphorylates the sites phosphorylated by MLCK (Thr-18, Ser-19), but the phosphatases affecting the physiologically important PKC sites on the MLC (Ser-1, Ser-2) or the MHC (Ser-1917) are unknown. There is growing
evidence however, of a role for phosphatases in controlling myosin function in platelets during the massive shape changes that occur during aggregation and secretion (Nakai et al., 1997; Toyoda et al., 1996). Interestingly, in these studies there was a greater association of PP2A with the cytoskeleton after stimulation than PP1, yet the literature suggests that PP1 plays the major role in myosin dephosphorylation. In light of these findings and our recent demonstration of the critical role of PP2A in membrane events during mast cell secretion (Ludowyke et al., 2000), we investigated the association of PP1 and PP2A with the rearranging actomyosin cytoskeleton during mast cell secretion.
MATERIALS AND METHODS

Cell Culture, Activation and Analysis of Secretion

Tissue culture supplies were obtained from Life Technologies, Inc. (Grand Island, NY, USA). Maintenance of monolayer cultures of RBL-2H3 cells was in RPMI with 10% fetal calf serum and carried out as described previously (Kawasugi et al., 1995; Ludowyke et al., 1989; Pfeiffer and Oliver, 1994). 1.5 x 10^6 cells/well in 6-well culture dishes or 1 x 10^5 cells/well in 8-well chamber slides (Becton Dickinson, Franklin Lakes, NJ, USA) were washed twice and activated at 37°C in Buffer A, consisting of 119 mM NaCl, 5 mM KCl, 5.6 mM dextrose, 0.4 mM MgCl_2 and 25 mM PIPES (1,4-piperazinediethane-sulphonic acid; pH 7.2). Buffer A was supplemented with 0.1% BSA and 1 mM CaCl_2 prior to use. For antigen activation, cells were primed overnight with 75 ng/ml DNP-specific IgE (Sigma, St Louis, MO, USA) and activated with 100 ng/ml of the specific antigen, DNP24-BSA (24 molecules of DNP conjugated with one molecule of BSA, hereafter referred to as DNP-BSA; Calbiochem-Novabiochem, La Jolla, CA, USA). Where required, cells were activated with 50 nM PMA or the combination of 50 nM PMA and 500 nM calcium ionophore A23187 (both from Sigma). Control samples were incubated with Buffer A alone. After the designated time, the reaction was stopped by placing the samples on ice; the supernatant was removed and an aliquot taken for assay of the release of β-hexosaminidase. The total cellular content of β-hexosaminidase was determined following lysis of unstimulated cultures and the activated release was expressed as a percentage of totals. The amount of β-hexosaminidase released was assayed using an absorbance assay with p-nitrophenyl-N-acetyl-β-D-glucosaminide (Sigma) as the substrate (Ludowyke et al., 2000). When isotopic labelling of cells was required, cells were labelled with 100 µCi/ml [32P] orthophosphoric acid (Amersham Pharmacia Biotech, Uppsala, Sweden) at 37°C for 2 hours in Buffer A with the addition of 4 mM glutamine, as previously described (Ludowyke et al., 1989; Ludowyke et al., 1996). The cells were washed three
times with Buffer A before activation (as above) or addition of inhibitors. When inhibitors were required, Okadaic acid (OA; potassium salt; Calbiochem-Novabiochem) was dissolved in Buffer A and jasplakinolide (Molecular Probes, Inc, Eugene, OR, USA) was dissolved in dimethyl sulfoxide. The final buffer concentration of dimethyl sulfoxide was no more than 0.2%, which was added to control cells and had no effects on secretion or morphology (Ludowyke et al., 1998). Cells were pre-incubated with the inhibitors in activation buffer containing 4 mM glutamine at 37°C for the designated times. Washing the cells free of excess inhibitors had no effect on the inhibitory capacity. Incubation of cells for 45 min or 3 hours in buffer alone had no effect on morphology or secretion.

**Fluorescence Microscopy**

After activation in an 8-well chamber slide as above, cells were fixed with 200 µl of a solution containing 3% v/v formaldehyde, 0.1% v/v gluteraldehyde and 0.1% Triton X-100 in phosphate-buffered saline (PBS) for 15 min at room temperature. The wells were washed in PBS and non-specific binding sites were blocked with 5% w/v skim milk in PBS for 30 min. The wells were incubated for 2 hours with specific rabbit polyclonal antibodies raised against peptides derived from the C termini of the catalytic subunits of PP1 or PP2A (as previously defined in (Sim et al., 1998)), or against non-muscle myosin (Biomedical Technologies Inc, Stoughton, MA, USA). After washing with PBS the wells were then incubated for 60 min with the FITC-labelled secondary antibodies (Amersham Pharmacia Biotech). F-actin was primarily detected using FITC-labelled phalloidin (Sigma) for 45 min. Alternatively, actin was detected using a monoclonal primary antibody for 2 hours followed by an FITC-labelled secondary antibody for 60 min (both from Amersham Pharmacia Biotech). A coverslip was then mounted on top of anti-quench reagent (90% glycerol, 10% v/v PBS, containing 1mg/ml p-phenylenediamine). Cells were viewed using an Olympus BX60 fluorescence microscope (Olympus, Japan) and Sensicam Imaging (Kelheim, Germany). Images were collected using a x60 oil objective directly into IP Lab Gel before final analysis in Adobe Photoshop. Under the
same conditions, using any of the primary or secondary antibodies alone, no signal was
detected (data not shown).

**Fluorescence intensity measurements**
Images of immunostained cells collected directly from the microscope as described above
were imported into NIH Image. For single whole cell analysis, a line was marked through the
cell, starting outside the periphery at one edge, extending through the nucleus, ending outside
the periphery at the opposite edge of the cell. For partial cell analysis, the line was marked
from outside the peripheral edge of the cell, 10 µm into the cell (the edge of the nucleus),
giving 90 point measurements. Up to three line measurements were taken from at least ten
cells from a number of different experiments, giving a total of 30 measurements. Means and
standard errors of the data points were used to plot the fluorescence intensity graphs. Intensity
graphs show symbol and error bar at every fifth data point. To compare between the control
and stimulated cells, ten points around the peak intensity of the peripheral band or lowest
points of the cytoplasmic clearance were compiled and compared with the equivalent points
from unstimulated cells or cells 30 min after activation. GraphPad Prism (GraphPad Software,
San Diego, CA, USA) was used to analyse the data using a non-parametric paired t-test
(Wilcoxon signed rank test, two-tailed, confidence interval 95%) giving the significance data
shown.

**Cell Lysis for Immunoprecipitation and Western Blotting**
After activation in a 6-well plate as above, 0.3 ml of ice-cold lysis buffer (Buffer B or C; see
below) was added and the cells were scraped immediately into microfuge tubes. The
immunoprecipitation of myosin was carried out as previously described, using Buffer B,
containing 250 mM NaCl, 100 mM Sodium Pyrophosphate, 100 mM Sodium Fluoride, 10
mM EGTA, 5 mM EDTA, 25 mM Tris-HCl (pH 8.5), 0.5% Nonidet P-40, 200 µM Pefabloc,
20 µg/ml leupeptin, pepstatin and aprotinin, 10 µM DNase and 10 µg/ml RNase (Ludowyke
et al., 1989; Ludowyke et al., 1996). Buffer C was used for immunoprecipitation of the
phosphatases and contained 100 mM NaCl, 50 mM Sodium Pyrophosphate, 50 mM Sodium Fluoride, 2 mM Sodium Orthovanadate, 20 mM HEPES (pH 8.8), 0.5% Nonidet P-40, 200 μM Pefabloc, 20 μg/ml leupeptin, pepstatin and aprotinin, 10 μM DNase and 10 μg/ml RNase (adapted from (Begum, 1995)). Monoclonal antibodies against the catalytic subunit of PP1 (Santa Cruz Biotechnology, Santa Cruz, CA, USA) or PP2A (Upstate Biotechnology, Lake Placid, NY, USA), or a polyclonal antibody against myosin (as above) were pre-bound for two hours to Protein A-Sepharose CL-4B (Amersham Pharmacia Biotech). Immunoprecipitation results achieved with the commercial anti-myosin antibody were the same as achieved using the well described anti-human platelet myosin antibody, kindly donated by Dr. Robert S. Adelstein, NIH, Bethesda, MD, USA (data not shown) and as previously described (Ludowyke et al., 1989). The cell lysates were rotated at 4°C for 12-16 hours with antibody-protein A-Sepharose beads. The lysate/bead slurry was subsequently added to ProbeQuant G-50 microcolumns (Amersham Pharmacia Biotech; pre-washed to remove G-50 resin) and centrifuged in screw capped tubes at 6,000 rpm for 20 seconds to remove unbound proteins. The beads were washed to remove non-specific proteins in lysis buffer, 50:50 lysis buffer:PBS and finally PBS. Beads were resuspended in 100 μl Laemmli sample buffer and heated at 85°C for 5 min to dissociate proteins from the beads.

**Western Blotting**

Immunoprecipitated proteins, solubilized in Laemmli sample buffer, were then separated on 10% SDS-polyacrylamide gels and transferred to polyvinylidene difluoride membranes at 400 mA for 2 hours. The membranes were washed and non-specific binding sites were blocked with 5% w/v skim milk in PBS. The membranes were incubated for 90 min with the polyclonal or monoclonal PP1 or PP2A primary antibodies or polyclonal myosin primary antibodies (as above) and 60 min with the horseradish peroxidase-labelled secondary antibodies (Amersham Pharmacia Biotech). The proteins were detected using an Enhanced Chemiluminescent detection kit (NEN Life Sciences Products, Boston, MA, USA) following
the manufacturer’s instructions. As a control, incubation of the cell lysate with protein-A-
sepharose beads alone, under the same conditions as above, showed no non-specific binding
of myosin, PP1 or PP2A. For quantitation, the bands on the Amersham x-ray film were
captured using a laser densitometer and analyzed using ImageQuant (Molecular Dynamics,
Sunnyvale, CA, USA). GraphPad Prism was used to analyse the densitometric data, using a
non-parametric paired t-test (two-tailed, confidence interval 95%) giving the significance data
shown.

One-Dimensional Isoelectric Focusing (IEF) and Tryptic Peptide Mapping

Immunoprecipitated myosin from $^{32}$P-labelled cells was separated by SDS-PAGE on either
12.5% gels for the light chain, or 5% gels for the heavy chain. Gels were stained with
Coomassie Blue (Bio-Rad, Hercules, California, USA), dried and subjected to
autoradiography. The areas containing the $^{32}$P-labelled myosin heavy or light chains were cut
from the gel and treated as described previously (Ludowyke et al., 1989). Briefly, gel slices
were digested overnight in a solution containing L-1-tosylamido-2-phenylethyl chloromethyl
ketone-trypsin in 50 mM NH$_4$HCO$_3$, centrifuged to remove gel slices, lyophilized and
resuspended in a final volume of 22 µl of 50 mM NH$_4$HCO$_3$. Tryptic peptide mapping was
conducted using a 0.5 mm-thick urea polyacrylamide one-dimensional IEF system as
previously described (Ludowyke et al., 1996; Moussavi et al., 1993). Briefly, the Biolytes
(Bio-Rad) used were pH 3-5, 4-6 and 5-8 (1:1:1). The peptides were separated by
electrophoresis at 25 W/1350 V at 10°C for 35 min with 1M NaOH as the cathode buffer and
1M H$_3$PO$_4$ as the anode buffer. The dried gels were then subjected to autoradiography and
analyzed as for Western blotting above.
RESULTS

Antigen-induced rearrangement of cytoskeletal proteins and phosphatases.

In unstimulated RBL-2H3 cells there are no actomyosin microfilaments evident, however actin and myosin are distributed throughout the cell and localized to small microvilli on the apical surface. Following the addition of antigen to IgE-primed mast cells, large ruffles are formed on the apical surface in which both actin and myosin become concentrated (Figure 1A; panels b and d). These changes are observed at a time prior to or coincident with the peak rate of granule mediator release (2.5 min post-stimulation for antigen) (Ludowyke et al., 1994a; Pfeiffer et al., 1985; Sahara et al., 1990; Spudich, 1994).

In unstimulated mast cells, PP1 and PP2A are broadly distributed throughout the cytoplasm and nucleus with a similar apical localization as actin and myosin (Figure 1B; panels a and c). Furthermore, as with myosin and actin, PP1 and PP2A become localized to the apical ruffles within 2.5 min after stimulation by antigen (Figure 1B; panels b and d). It is also evident that PP1 and PP2A are in the nucleus, but the levels do not change following activation (L. Scurr, W. Sewell, R. Ludowyke; unpublished observations).

In addition to the formation of these surface ruffles, there are many other changes in the morphology and cytoskeletal structure of mast cells after stimulation. The cells decrease in height by 50% (Ludowyke et al., 1994a; Ludowyke et al., 1994b) and spread out on the substrate, increasing their surface area (Ludowyke et al., 1994b; Oliver et al., 1988). As shown in Figure 2A, these changes correspond to considerable cytoskeletal rearrangements on the basal surface of the cells. Within 2.5 min after addition of antigen, a thick band of actin and myosin has formed at the periphery of the spreading cells, with a corresponding clearance of these proteins from the cytoplasm between the nucleus and cell periphery (Figure 2A; panels b and e). By 30 min post stimulation however, myosin and actin have returned to a distribution similar to that seen in unstimulated cells, although the actin plaques are still evident (Figure 2A; panels c and f).
Figure 2B documents a similar pattern of redistribution of PP1 and PP2A. Within 2.5 min of the addition of antigen, both PP1 and PP2A are cleared from the cytoplasm of the spreading cell and are concentrated in a thick band around the periphery of the cell (Figure 2B; panels b and e). This band can be seen forming within one minute after addition of antigen, indicating that these changes do in fact precede the peak rate of granule secretion (data not shown). However, within 30 min of stimulation, both PP1 and PP2A have returned to their cytoplasmic locations, with the pattern of distribution being similar to that found in unstimulated cells (Figure 2B; panels c and f).

In order to quantitate the changes in these cells, the fluorescence intensity was analysed. Initially, cross-sections from a single cell were taken, clearly defining the broad band of protein at the periphery of the cell, 2.5 min after activation (Figure 3A; panel b). The graphs also suggest that by 30 min post stimulation, the pattern of distribution has reverted to that of unstimulated cells (Figure 3A; panels c and a). To specifically analyse the band at the cell periphery and the clearance from the cytoplasm, fluorescence intensity data was compiled from a point immediately outside the periphery of the cell to the edge of the nucleus in both unstimulated and stimulated cells. These data indicate that antigen stimulation induces a significant (P<0.002), 2-fold increase in fluorescence intensity in the band at the cell periphery for PP1, PP2A and myosin (Figure 3B). There is a corresponding significant (P<0.002) 3-fold decrease in intensity approximately midway between the nucleus and cell periphery for PP1 and PP2A and a 5-fold decrease in myosin immunostaining (Figure 3B). These data also confirm that by 30 min post stimulation, the distribution of PP1, PP2A and myosin have returned to that of unstimulated cells. Actin was not analysed as the staining was more punctate and there was no nuclear staining to orient the line measurements.

PMA plus calcium ionophore A23187-induced rearrangement of the cytoskeleton and phosphatases.
PMA plus A23187 induced secretion with a slower onset and a delayed peak rate of secretion; 4-6 min compared with 2.5 min for antigen (Ludowyke et al., 2000). As seen with antigen stimulation a peripheral accumulation of PP1 and PP2A was also observed at the peak of PMA plus A23187-stimulated secretion (Figure 4A; panels b and e). The clearance of PP1 and PP2A from the cytoplasm between the nucleus and cell periphery is slightly more pronounced than seen with antigen (Figure 4B; 5-6-fold decrease in fluorescence intensity, P<0.002). Interestingly, unlike antigen stimulation, the distribution of PP1 and PP2A does not return to unstimulated levels 30 min after addition of PMA plus A23187 (Figure 4A; panels c and f). At this time, the clearance from the cytoplasm remains significantly different from unstimulated cells (Figure 4B; 3-4-fold, P<0.002). These results show that two separate secretagogues with different initial modes of action and time-course, both induce a similar redistribution of PP1 and PP2A at the time of peak granule secretion. A similar rearrangement of actin and myosin at these time points was observed (data not shown).

The addition of PMA alone to RBL-2H3 cells does not induce secretion, yet it does induce many of the intracellular signals generated by antigen, such as activation of PKC and a redistribution of actin and myosin (Choi et al., 1994; Ludowyke et al., 1994b; Ludowyke et al., 1996). PMA alone induces a similar rearrangement of PP1 and PP2A as that seen with PMA plus A23187 (Figure 4A), suggesting that these changes do not require an increase in intracellular calcium (data not shown).

**Co-Immunoprecipitation of Myosin, PP1 and PP2A**

Having established that during the peak rate of granule mediator release, PP1 and PP2A are localized in areas where myosin and actin are also concentrated, we sought to determine if there was an association between these proteins. Myosin, PP1 and PP2A were immunoprecipitated from unstimulated and antigen stimulated RBL-2H3 cells as described in MATERIALS AND METHODS (Ludowyke et al., 1989; Ludowyke et al., 1996). The
myosin immunocomplex was probed by Western immunoblotting for the presence of PP1c and PP2Ac. In reciprocal analyses, PP1c or PP2Ac immunocomplexes were probed for the presence of myosin. As these reciprocal immunoprecipitations investigated the association of myosin with the phosphatases, the densitometric analyses of the Western blots were combined. Whether myosin or the phosphatases were immunoprecipitated, relative to unstimulated cells, there was a significant increase in the association of myosin with both PP1c and PP2Ac at the peak rate of secretion (Figure 5). There was a 1.78 fold $\pm$ 0.25 (n=5; $P<0.05$) increase in the association of myosin with PP1c and a 1.36 fold $\pm$ 0.04 (n=6; $P<0.05$) increase in the association of myosin with PP2Ac. Remarkably, by 5 min post-stimulation the amount of myosin associated with both PP1c and PP2Ac had returned to levels similar to that in unstimulated cells (1.05 $\pm$ 0.11 and 1.09 $\pm$ 0.14 fold respectively; Figure 5). The level of recovery of PP1, PP2A and myosin from their respective immunoprecipitates was the same from the control and at each time point (data not shown).

**Inhibition of mast cell secretion by inhibitors of actin assembly and phosphatase activity.**

To further investigate the link between cytoskeletal rearrangements, PP1, PP2A and secretion, we used jasplakinolide, an inhibitor of actin assembly and Okadaic acid (OA), an inhibitor of phosphatase activity. Jasplakinolide has been shown to bind to actin, inhibiting polymerization (Bubb et al., 1994). We have recently shown that jasplakinolide can inhibit insulin secretion from RINm5F cells (Wilson et al., 2001). We have found here that pre-incubation of RBL-2H3 cells for 3hrs with 5 $\mu$M jasplakinolide, inhibits the antigen-induced secretion of $\beta$-hexosaminidase by 73% $\pm$ 9.4%. Under these conditions, jasplakinolide does not affect the association of either PP1 or PP2A with myosin in unstimulated cells. Furthermore, although jasplakinolide inhibited secretion, there was no effect on the increased association of myosin with PP1 and PP2A, 2.5 min after antigen stimulation (Figure 6A). In addition, the association of actin with myosin does not appear to be affected by jasplakinolide.
or to change over this time period (Figure 6A). Nevertheless, in antigen stimulated cells, jasplakinolide prevented the clearance of actin, PP1 and PP2A from the cytoplasm and the formation of the characteristic band at the periphery of the cells (Figure 7; panels a, c, and e). However, these cytoskeletal changes are seen in cells preincubated with buffer for 3 hours and subsequently stimulated with antigen (Figure 7; panels b, d, and f). Control cells incubated with jasplakinolide, without subsequent addition of antigen were similar to those depicted in the presence of antigen (data not shown). Jasplakinolide induced some of the cells to become round, however there were few other effects of jasplakinolide upon the distribution of actin, whether investigated using a monoclonal anti-actin antibody or phalloidin (data not shown).

Surprisingly, there was a very clear alteration in the distribution of myosin following the addition of jasplakinolide. Jasplakinolide induced myosin to form short, rod-like structures throughout the cell (highlighted in a magnified view, Figure 7; panel g (ii)). These structures remained after antigen stimulation and there was no clearance from the cytoplasm or formation of the peripheral band. Interestingly, jasplakinolide did not affect the association of myosin with PP1, PP2A or actin, before or after stimulation (Figure 6A). These data suggest that antigen-mediated PP1 and PP2A redistribution is dependent upon an intact actomyosin microfilament system, which is also required for secretion in these cells.

Addition of the PP1 and PP2A inhibitor OA to RBL-2H3 cells inhibited secretion (Kitani et al., 1996; Ludowyke et al., 1998), however we have shown that the addition of 1 µM OA for 45 min only inhibits the activity of PP2A (Ludowyke et al., 2000). In the present experiments using the same conditions, OA inhibited the antigen-induced secretion of β-hexosaminidase by 74% ± 4%, a similar amount to that of jasplakinolide.

Analysis of myosin immunoprecipitates revealed that the addition of OA inhibited the association of PP1 and PP2A with myosin in unstimulated cells by approximately 50% (data not shown). Furthermore, where previously the association of PP2A with myosin increased
2.5 min after antigen stimulation (Figure 5B), OA pre-treatment led to a decreased association at this time point (Figure 6B). PP1 was not detected in association with myosin after OA pre-treatment, although it was clearly evident after immunoblotting the unbound proteins (data not shown). There was no apparent change in the amount of actin associated with myosin over this time course (Figure 6B). Consistent with these findings, immunofluorescence images showed that OA clearly disrupts the antigen-stimulated rearrangement of actin and myosin. Following pre-incubation with OA, the cells became round and the addition of antigen did not induce any cytoskeletal alterations. That is, there was no formation of the characteristic band surrounding the periphery of the cells or clearance of actin, myosin, PP1 and PP2A from the cytoplasm (Figure 8; panels a, c, e and g). However, these cytoskeletal changes are seen in cells pre-incubated with buffer and subsequently stimulated with antigen (Figure 8; panels b, d, f and h). Control cells pre-incubated with OA without subsequent addition of antigen were similar to those depicted here in the presence of antigen (data not shown).

**Inhibition of phosphatase activity leads to increased MHC and MLC phosphorylation at PKC-specific sites**

Having previously established that pre-incubation of RBL-2H3 cells with OA (1 µM for 45 min) inhibited the activity of PP2A and not that of PP1 (Ludowyke *et al.*, 2000), our present data implied that the cytoskeletal changes we observed here may also be regulated more by PP2A than PP1. To further investigate this, we added 1 µM OA to cells that had been pre-labelled with $^{32}$P to label intracellular pools of ATP and then immunoprecipitated myosin as described in MATERIALS AND METHODS. Both the 200 kDa MHC and the 20 kDa MLC are phosphorylated in unstimulated cells and the level of phosphorylation increases when the cells are stimulated by antigen or PMA (Figure 9A and (Choi *et al.*, 1994; Ludowyke *et al.*, 1989; Ludowyke *et al.*, 1996)). We demonstrate here that the addition of OA also leads to a time-dependent increase in the phosphorylation of the MHC’s and MLC’s
(Figure 9A). The MHC’s and MLC’s were excised from the gels, trypsinized, lyophilized and the tryptic peptides separated by one-dimensional IEF as previously described (Ludowyke et al., 1996). The peptide map of the MHC shows a number of phosphopeptides in control cells (Figure 9B). The addition of OA leads to the time-dependent generation of another phosphopeptide separating towards the cathode. Although increased phosphorylation is seen in other peptides, this particular phosphopeptide matches the MHC phosphopeptide generated following the addition of PMA to Jurkat cells and antigen to RBL-2H3 cells (Ludowyke et al., 1989; Moussavi et al., 1993) and is known to contain the PKC phosphorylation site Ser-1917 (Conti et al., 1991).

The peptide map of the MLC shows a single phosphopeptide in control cells, previously established as the Ser-19 site that is phosphorylated by MLCK. As we have previously demonstrated, the addition of antigen or PMA leads to the phosphorylation of another site, Ser-1 or Ser-2, that is known to be phosphorylated by PKC (Ludowyke et al., 1989; Ludowyke et al., 1996). The addition of OA induced a time-dependent increase in phosphorylation of these PKC-specific sites (Figure 9C). Although in some experiments there was a small increase in the phosphorylation at the MLCK and Di-MLCK (Thr18-Ser19) sites, the predominant effect of the addition of OA was an increased phosphorylation at the PKC sites. The two MLC peptides attributed to phosphorylation by PKC (PKC and PKC') both correspond to Ser-1 or Ser-2, but the upper band is generated by further tryptic cleavage of the arginine residue from the sequence Ser-Ser-Lys-Arg (Moussavi et al., 1993; Nakabayashi et al., 1991).

Therefore the addition of OA to RBL-2H3 cells time-dependently inhibits the activity of PP2A and leads to increased phosphorylation of the MHC and MLC at sites known to be phosphorylated specifically by PKC.
DISCUSSION

Regulated exocytosis requires the remodelling of the cortical actomyosin barrier, allowing granules access to their fusion sites to release their contents into the extracellular space. In mast cells, the kinetics of this remodelling correlates with the increased phosphorylation of myosin on both its heavy and light chains, at serine and threonine sites. Some of the kinases responsible for this increased phosphorylation are MLCK, PKC and CaM kinase II. However, increased phosphorylation can also be induced by the altered activity of a phosphatase, yet there is little information about the serine/threonine phosphatases that regulate myosin function during exocytosis in any cell. Here we demonstrate for the first time, that following mast cell stimulation, both the major serine/threonine phosphatases, PP1 and PP2A associate with myosin and play a major role in regulating the cytoskeletal remodelling.

Our data clearly demonstrates that following activation, PP1 and PP2A are relocated from a diffuse distribution throughout the cytoplasm to regions of the cell where actin and myosin have become concentrated. Recent work has shown that the apical ruffles formed in activated RBL-2H3 cells also contain PLCγ1 (Barker et al., 1998), suggesting that they are a locus of high cellular signalling activity during the secretory process. The dynamic nature of these changes is demonstrated in the time course of the redistribution of PP1 and PP2A following stimulation which coincides or indeed precedes the time of peak release of granule mediators (Ludowyke et al., 2000). This correlation is also apparent in the slower, more prolonged response to PMA plus A23187, due to the prolonged activation of PKC by phorbol esters. As actin and myosin are similarly redistributed, it is feasible that PP1 and PP2A contribute to these cytoskeletal rearrangements.

The interaction between actin and myosin is primarily regulated by phosphorylation. It is clear that in vitro or in living cells, PP1 can associate with and dephosphorylate the myosin light chain and affect actin microfilaments (Chisholm and Cohen, 1988; Fernandez et al.,...
In unstimulated RBL-2H3 cells, both PP1 and PP2A associate with myosin and following stimulation this level of association is significantly increased. Remarkably, this increased association is transient, coinciding with the peak rate of secretion, and as the rate of secretion diminished, so too did the association of the phosphatases with myosin. This suggests there is an active mechanism for attachment and removal of the phosphatases from the myosin complex. In our previous study we described the transient translocation of phosphatases to the plasma membrane of RBL-2H3 cells, again peaking at 2.5 min (Ludowyke et al., 2000). However, in that instance only PP2A was translocated to the plasma membrane; PP1 was not. Therefore, whilst both PP1 and PP2A contribute to the cytoskeletal changes required for secretion, only PP2A is involved in the secretory events at the plasma membrane.

Jasplakinolide binds to and stabilizes F-actin, limiting the amount of monomeric actin available for remodelling of stress fibres (Bubb et al., 2000). We have recently demonstrated that jasplakinolide inhibits insulin secretion from RINm5F cells (Wilson et al., 2001). In the present study jasplakinolide inhibited secretion from mast cells and prevented the cytoskeletal rearrangement of actin, PP1 and PP2A seen after antigen stimulation. Surprisingly, jasplakinolide also had a dramatic effect on myosin, inducing the formation of short rods, spread throughout the cytoplasm. This distribution was unaffected by antigen stimulation. To our knowledge, this is the first time such an effect of jasplakinolide on myosin has been described. Interestingly, jasplakinolide did not affect the association of myosin with PP1, PP2A or actin, before or after stimulation. Therefore, inhibition of secretion by jasplakinolide can be interpreted as evidence of the requirement for a dynamic actomyosin interaction in the secretory process.

Addition of OA to RBL-2H3 cells induced changes in the association of both PP1 and PP2A with myosin and inhibited the activated cytoskeletal changes. However in RBL-2H3 cells, under these conditions, OA only inhibits the activity of PP2A, suggesting that PP2A has
the predominant role in these cytoskeletal changes and may also effect the association of PP1 with myosin. PP2A is the major phosphatase acting on caldesmon, a protein that blocks the interaction of actin with myosin (Ferrigno et al., 1993). Furthermore, in fibroblasts, dephosphorylation of the intermediate filament protein vimentin has been shown to be controlled by PP2A and not PP1 (Turowski et al., 1999) and the addition of OA to RBL-2H3 cells leads to increased vimentin phosphorylation (Ludowyke et al., 1998). Therefore it is likely that dynamic changes in the phosphorylation state of several cytoskeletal proteins is regulated by PP2A (Sontag, 2001). Our present finding that PP2A associates with myosin during mast cell secretion was however, surprising, because only PP1 has been implicated in the regulation of myosin through MLC phosphorylation (Chisholm and Cohen, 1988; Hartshorne et al., 1998). However, recent studies in Dictyostelium have implicated PP2A as the regulatory phosphatase controlling myosin heavy chain phosphorylation at sites that regulate myosin filament assembly (Murphy and Egelhoff, 1999). In RBL-2H3 cells, others and we have shown that activation induces the phosphorylation of the MHC by PKC and Cam Kinase II. These sites are close to the carboxyl terminus at the end of the coiled-coil region and are believed to contribute to the rearrangement of the actomyosin cytoskeleton that occurs during the secretory process (Buxton and Adelstein, 2000; Ludowyke et al., 1989). In unstimulated RBL-2H3 cells the MLC is phosphorylated at a specific site (Ser-19) that is regulated by MLCK and PP1, but phosphorylation at this site does not change following activation. However, activation leads to a significant increased phosphorylation at specific sites (Ser-1/Ser-2) that are the major physiological sites for PKC (Choi et al., 1994; Ludowyke et al., 1989; Ludowyke et al., 1996). The phosphatase that regulates phosphorylation at these PKC sites is unknown, but our present data implies that the major phosphatase involved is PP2A. The addition of OA under conditions that inhibit only PP2A and not PP1, induces little alteration in the phosphorylation of the MLC sites regulated by MLCK and PP1, but a significant increase in the phosphorylation of sites regulated by PKC.
Therefore our findings suggest that PKC and PP2A together may be involved in regulating the phosphorylation of the MHC at Ser-1917 and the MLC at Ser-1/Ser-2 that occurs during mast cell secretion.

Although there is no evidence for OA mediated activation of PKC, the increased phosphorylation at PKC-specific sites on myosin may occur in a number of ways. The inhibition of PP2A may lead to the stimulation of an upstream activator of PKC or the removal of an inhibitory binding protein. Alternatively, the unstimulated activity of PKC for myosin may always be high, yet the level of MHC or MLC phosphorylation is kept low by a much greater activity of PP2A. Thus inhibiting the activity of PP2A leads to increased phosphorylation due to the high level of unstimulated PKC activity. Whilst these are important questions, the major impact of the results presented here is that PP2A is identified as playing an important role in regulating the phosphorylation and thus the function of myosin during mast cell secretion.

In the larger context, a question arises as to what functional role these changes have in the mast cell exocytotic process. Notably, in the area of the cytoplasm between the nucleus and the cell periphery, there was a significant decrease in the levels of PP1 and PP2A (3.2 fold) and myosin (5 fold). This “clearance” of the actomyosin cytoskeleton from this region may be an important mechanism to allow granules easier access to the plasma membrane. We propose that the formation of the peripheral band pulls at the cortical barrier from one direction (towards the periphery), and the formation of the apical ruffles pulls the barrier back towards the centre of the cell or at least in an opposing direction, thus “stretching” the cortical web in the region between the nucleus and cell periphery. As the cells spread out by about 50% (Ludowyke et al., 1994a; Ludowyke et al., 1994b), this region, which has a smaller distance between the apical and basal surfaces, also has a diminished concentration of actomyosin, which may allow the granules easier access to the plasma membrane, without requiring a major disassembly of the cortical barrier.
In summary, stimulation of RBL-2H3 cells induces a transient association of the serine/threonine phosphatases PP1 and PP2A with myosin, which coincides with the peak rate of granule mediator release. This association is likely to be linked with the transient but significant clearance of actomyosin from the cytoplasm between the cell periphery and the nucleus, into ruffles on the apical surface and a band at the periphery of the cell. This remodelling coincides with or precedes the peak rate of mediator release. Inhibitors of either phosphatase activity or actin assembly inhibit both exocytosis and this cytoskeletal remodelling and give evidence of the interaction of PKC and PP2A in regulating myosin phosphorylation. Overall, our studies indicate that exocytosis does require a dynamic actomyosin microfilament system and that PP1 and PP2A play a stimulatory role in these events.
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ABBREVIATIONS

The abbreviations used are: DNP-BSA, antigen containing 24 molecules of dinitrophenol (DNP) conjugated with 1 molecule of bovine serum albumin (BSA); FITC, fluorescein isothiocyanate; IEF, isoelectric focusing; MHC, myosin heavy chain; MLC, myosin light chain; MLCK, myosin light chain kinase; OA, Okadaic acid; PBS, phosphate buffered saline; PKC, protein kinase C; PMA, phorbol 12-myristate 13-acetate; PP1 or PP2A, protein phosphatase type 1 or type 2A; RBL-2H3, rat basophilic leukemia cell line, subclone 2H3.
REFERENCES


FIGURE LEGENDS

Figure 1 - Redistribution of actin, myosin, PP1 and PP2A to apical ruffles after antigen stimulation.
Adherent monolayers of RBL-2H3 cells that had been incubated overnight with DNP-specific IgE were incubated in buffer alone for 2.5 min (Unstimulated, panels a, c) or with antigen (100 ng/ml DNP-BSA) for 2.5 min (panels b, d), fixed and proteins on the apical surface detected as detailed under MATERIALS AND METHODS. Actin (Figure 1A, panels a, b) was detected using phalloidin-FITC and myosin (Figure 1A, panels c, d), PP1c (Figure 1B, panels a, b) and PP2Ac (Figure 1B, panels c, d) using specific polyclonal antibodies and FITC-labelled secondary antibodies. The images focus on the apical surface of the cells and are representative of those seen in at least five separate experiments. Scale bar is 5 µm.

Figure 2 - Transient redistribution of actin, myosin, PP1 and PP2A to the periphery of the spreading cell after antigen stimulation.
Adherent monolayers of RBL-2H3 cells that had been incubated overnight with DNP-specific IgE were incubated in buffer alone for 2.5 min (Unstimulated, panels a, d), or with antigen (100 ng/ml DNP-BSA) for 2.5 min (panels b, e) or 30 min (panels c, f). The cells were then fixed and proteins on the basal cell surface detected as detailed under MATERIALS AND METHODS. Actin (Figure 2A, panels a-c) was detected using phalloidin-FITC and myosin (Figure 2A, panels d-f), PP1c (Figure 2B, panels a-c) and PP2Ac (Figure 2B, panels d-f) using specific polyclonal antibodies and FITC-labelled secondary antibodies. Arrows highlight regions where protein has cleared from the cytoplasm and arrowheads highlight the band of protein around the peripheral edge. The lines across the cells immunostained with PP1c were used to generate the fluorescence intensity graphs of Figure 3A. The images focus on the basal surface of the cells and are representative of those seen in at least five separate experiments. Scale bar is 5 µm.
Figure 3- Fluorescence intensity graphs of antigen stimulated cells.

NIH Image was used to generate fluorescence intensity data from immunostained cells as depicted in Figure 2. The line across the cells immunostained with PP1c (Figure 2B, panels a-c) was used to generate the fluorescence intensity graphs from unstimulated cells and those stimulated with antigen for 2.5 and 30 min (Figure 3A, panels a, b and c respectively). Figure 3B focuses on the area between the edge of the cell and the beginning of the nucleus. Three line measurements were taken from at least ten cells and the data compiled as described under MATERIALS AND METHODS from cells immunostained with PP1, PP2A and myosin. The data (mean ± SE) from unstimulated cells and those stimulated with antigen for 2.5 and 30 min are plotted together for comparison. Intensity graphs show symbol and error bar at every fifth data point. Numerical data above the graphs indicates the difference between unstimulated and 2.5 min antigen stimulated cells at the peripheral band (fold increase) or the clearance (fold decrease). The significance of this difference is indicated (** P<0.002).

Figure 4 - Relocation of PP1 and PP2A to the periphery of the spreading cell after PMA plus A23187 stimulation.

Adherent monolayers of RBL-2H3 cells were incubated in buffer alone for 6 min (a, d) or with 50 nM PMA plus 500 nM A23187 for 6 min (panels b, e) or 30 min (panels c, f), fixed and proteins detected as detailed under MATERIALS AND METHODS. PP1c (panels a-c) and PP2Ac (panels d-f) were detected using specific polyclonal antibodies and FITC-labelled secondary antibodies. Arrowheads highlight the band of protein around the peripheral edges of the cells and arrows highlight regions where protein has cleared from the cytoplasm. The images focus on the basal surface of the cells and are representative of those seen in at least three separate experiments. Scale bar is 5 µm. A fluorescence intensity graph of compiled line measurements from the edge of the cell to the beginning of the nucleus as described in the
legend to Figure 3 above, is depicted (Figure 4B). The data (mean ± SE) from unstimulated cells and those stimulated with PMA plus A23187 for 6 and 30 min are plotted together for comparison. Numerical data above the graphs indicates the difference in the clearance (fold decrease) between unstimulated and 6 min PMA plus A23187 stimulated cells. The significance of this difference is indicated (** P<0.002).

**Figure 5 - Coimmunoprecipitation of myosin and PP1/PP2A after antigen stimulation.**

Adherent monolayers of RBL-2H3 cells that had been incubated overnight with DNP-specific IgE were incubated in buffer alone or activated with antigen (100 ng/ml DNP-BSA) for 2.5 or 5 min. Cells were lysed and PP1c, PP2Ac or myosin immunoprecipitated using specific antibodies as detailed under MATERIALS AND METHODS. PP1c and PP2Ac immunoprecipitates were probed for myosin by Western blotting and myosin immunoprecipitates were probed for PP1c and PP2Ac as above. The images are representative of those seen in two to three separate experiments.

**Figure 6 - Effect of Jasplakinolide or Okadaic acid on coimmunoprecipitation of myosin and PP1/PP2A.**

Adherent monolayers of RBL-2H3 cells that had been incubated overnight with DNP-specific IgE were incubated in a buffer containing either 5 µM Jasplakinolide (Figure 6A) for 3 hours or 1 µM OA (Figure 6B) for 45 min. Cells were subsequently left in buffer alone or activated with antigen (100 ng/ml DNP-BSA) for 2.5 min, lysed and myosin immunoprecipitated using the anti-myosin polyclonal antibody as detailed under MATERIALS AND METHODS. Myosin immunoprecipitates were probed for PP1c, PP2Ac and actin by Western blotting. The images are representative of those seen in two separate experiments.
Figure 7 - Effect of Jasplakinolide on relocation of actin, PP1, PP2A and myosin after antigen stimulation.

Adherent monolayers of RBL-2H3 cells that had been incubated overnight with DNP-specific IgE were incubated in a buffer containing 5 µM Jasplakinolide for 3 hours and activated with antigen (100 ng/ml DNP-BSA) for 2.5 min (panels a, c, e and g). A separate set of cells was left for 3 hours in buffer alone as a control and activated with antigen (100 ng/ml DNP-BSA) for 2.5 min (panels b, d, f and h). All cells were fixed and proteins were detected as detailed under MATERIALS AND METHODS. Actin (panels a, b) was detected using an anti-actin monoclonal antibody and FITC-labelled secondary antibody, PP1c (panels c, d), PP2Ac (panels e, f) and myosin (panels g, h) were detected using specific polyclonal antibodies and FITC-labelled secondary antibodies. Panel g (ii) is an enlarged view of Jasplakinolide treated unstimulated cells highlighting the short rods of myosin. The images focus on the basal surface of the cells and are representative of those seen in at least three separate experiments. Scale bar is 5 µm.

Figure 8 - Effect of Okadaic acid on relocation of actin, myosin, PP1 and PP2A after antigen stimulation.

Adherent monolayers of RBL-2H3 cells that had been incubated overnight with DNP-specific IgE were incubated in a buffer containing 1 µM OA for 45 min and activated with antigen (100 ng/ml DNP-BSA) for 2.5 min (panels a, c, e and g). A separate set of cells was left for 45 min in buffer alone as a control and activated with antigen (100 ng/ml DNP-BSA) for 2.5 min (panels b, d, f and h). All cells were fixed and proteins were detected as detailed under MATERIALS AND METHODS. Actin (panels a, b) was detected using an anti-actin monoclonal antibody and FITC-labelled secondary antibody, myosin (panels c, d), PP1c (panels e, f) and PP2Ac (panels g, h) were detected using specific polyclonal antibodies and
FITC-labelled secondary antibodies. The images focus on the basal surface of the cells and are representative of those seen in at least three separate experiments. Scale bar is 5 µm.

**Figure 9 - Phosphopeptide Mapping of Myosin Heavy and Light Chains**

Adherent monolayers of RBL-2H3 cells that had been incubated for 2 hours with 100 µCi/ml $^{[32P]}$ orthophosphoric acid were washed, then incubated in a buffer containing 1 µM OA for 15, 30 or 45 min, or activated with antigen (100 ng/ml DNP-BSA) or PMA (50nM) for 10 min. Cells were lysed and myosin immunoprecipitated using specific antibodies, before being separated on 5% or 12.5% SDS-PAGE and subjected to autoradiography as detailed under MATERIALS AND METHODS. Panel A shows an autoradiogram of a 12.5% gel depicting the changes in phosphorylation of the myosin heavy and light chains. The myosin heavy and light chains were then excised from the gel, digested with trypsin and the peptides separated on a one-dimensional IEF gel as detailed under MATERIALS AND METHODS. Panels B and C show the heavy chain and light chain peptide maps respectively. Direction of the IEF on both peptide maps is toward the anode which is at the top of Panels B and C. The peptide labelled PKC in the MHC phosphopeptide map corresponds to the peptide in platelet myosin phosphorylated *in vitro* by PKC (Moussavi *et al.*, 1993). The two MLC peptides that were phosphorylated by PKC (PKC and PKC') correspond to Ser-1 or Ser-2 and are generated by different tryptic cleavage (Moussavi *et al.*, 1993). MLCK forms peptides monophosphorylated at Ser-19 and diphosphorylated at Thr-18 and Ser-19. The amount of myosin immunoprecipitated was checked by Coomassie blue staining and confirmed there was little difference in loading between samples. The images shown are representative of 4 separate experiments.
FIGURE 3

3A

- a - Unstimulated
- b - Antigen, 2.5 min
- c - Antigen, 30 min

3B

- PP1
- Unstimulated
- Antigen, 2.5 min
- Antigen, 30 min
- 2.2**
- 3.2**

- PP2A
- Unstimulated
- Antigen, 2.5 min
- Antigen, 30 min
- 2.1**
- 3.2**

- Myosin
- Unstimulated
- Antigen, 2.5 min
- Antigen, 30 min
- 1.8**
- 5.0**
FIGURE 4

4A

Unstimulated

6 min

30 min

4B

Unstimulated

PMA + A23187, 6 min

PMA + A23187, 30 min

Fluorescence Intensity

Distance (µm)

5.0**

6.4**
FIGURE 5

A - PP1c/Myosin Co-immunoprecipitation

B - PP2Ac/Myosin Co-immunoprecipitation

FIGURE 6
FIGURE 7

3 hour Jas: 2.5 min Antigen

ACTIN

PP1c

PP2Ac

MYOSIN
FIGURE 8

45min OA:
2.5 min Antigen

45min Buffer:
2.5 min Antigen

ACTIN

MYOSIN

PP1c

PP2Ac
FIGURE 9

9A - Autoradiogram of Immunoprecipitated Myosin

200kDa Myosin Heavy Chain

1 - Control
2 - 15 min OA
3 - 30 min OA
4 - 45 min OA
5 - 10 min DNP/BSA
6 - 10 min PMA

20kDa Myosin Light Chains

9B - Myosin Heavy Chain Peptide Map

9C - Myosin Light Chain Peptide Map

PKC

PKC'

Di-MLCK

MLCK

PKC

Cont  OA  OA  OA  DNP/BSA  PMA
15min 30min 45min 10min 10min