Coronins are a conserved family of WD repeat-containing, actin-binding proteins that regulate cell motility in a variety of model organisms. Our results show that Coronin 1B is a ubiquitously expressed member of the mammalian Coronin gene family that co-localizes with the Arp2/3 complex at the leading edge of fibroblasts, and co-immunoprecipitates with this complex. Pharmacological experiments show that the interaction between Coronin 1B and the Arp2/3 complex is regulated by protein kinase C (PKC) phosphorylation. Coronin 1B is phosphorylated by PKC both in vitro and in vivo. Using tryptic peptide mapping and mutagenesis, we have identified serine 2 (Ser-2) on Coronin 1B as the major residue phosphorylated by PKC in vitro. Rat fibroblasts expressing the Coronin 1B S2A mutant show enhanced ruffling in response to phorbol 12-myristate 13-acetate (PMA) and increased speed in single cell tracking assays. Cells expressing the Coronin 1B S2D mutant have attenuated PMA-induced ruffling and slower cell speed. Expression of the S2A mutant partially protects cells from the inhibitory effects of PMA on cell speed, whereas expression of the S2D mutant renders cells hypersensitive to its effects. These data demonstrate that Coronin 1B regulates leading edge dynamics and cell motility in fibroblasts, and that its ability to control motility and interactions with the Arp2/3 complex are regulated by PKC phosphorylation at Ser-2. Furthermore, Coronin 1B phosphorylation is responsible for a significant fraction of the effects of PMA on fibroblast motility.

The dynamic re-organization of the actin cytoskeleton is required for many processes including migration, endocytosis, and intracellular junction formation. Coronins are a conserved family of WD repeat-containing, actin-binding proteins that regulate migration and other actin-dependent processes in model organisms. Mammalian genomes contain at least six Coronin genes, however, little is known about the molecular function or regulation of Coronins in mammalian cell motility. The founding member of the Coronin protein family was discovered in Dictyostelium as an F-actin-binding protein present in a contracted myosin-actin preparation (1). Subsequent gene knock-out studies showed that this protein was involved in cell motility and cytokinesis (2). Dictyostelium cells deficient in Coronin move at about half the rate of the wild-type controls and have a high frequency of failed cytokinasis. The protein is localized at the leading edge of motile cells. In addition to motility defects, Coronin null cells also have strong defects in fluid phase endocytosis (3).

Yeast also has a Coronin protein that binds to F-actin in vitro and localizes in vivo to dynamic actin structures such as actin patches (4, 5). Unlike Dictyostelium, disruption of the single Coronin gene (CRN1) in yeast does not lead to gross defects in growth or cytoskeletal organization. However, when a crn1 mutation is combined with a mutation in the gene encoding Coflin (cof1–22), a protein involved in actin filament turnover, synthetic defects in growth and cytoskeletal organization are observed (5). Overexpression of Coronin in yeast induces the formation of aberrant actin loops and causes lethality (6). Purified yeast Coronin binds directly to the Arp2/3 complex via its coiled-coil domain and inhibits Arp2/3-mediated actin nucleation in vitro. In support of these in vitro studies, mutations in the CRN1 gene genetically synergize with mutations in the subunits of the Arp2/3 complex.

Coronin has also been studied in other model organisms. In Drosophila, mutations in Coronin cause disruptions in the actin cytoskeleton of the embryonic imaginal disks and an early pupal lethal phenotype, indicating that this gene is essential for morphogenesis in this organism (7). In Xenopus, Coronin localizes to the periphery of fibroblastic cells and, like Dictyostelium and yeast Coronins, co-sediments with F-actin in vitro (8). Also, the coiled-coil region at the C terminus of Xenopus Coronin has been shown to mediate oligomerization (9). Expression of a putative dominant negative construct containing the five canonical WD repeats from Xenopus Coronin leads to defects in cell spreading and Rac-induced lamellipodial formation (8).

Several mammalian members of the Coronin family have been described (10). For clarity, we have adopted the Human Genome Organization (HUGO) nomenclature for mammalian Coronin genes (www.gene.ucl.ac.uk/nomenclature/genefamily/coron.html). Coronin 1A (also known as p57 or Coronin-1) was the first Coronin to be identified in mammals and is highly expressed in cells of the hematopoietic lineage as well as parts of the nervous system (11). This protein is a substrate for PKC both in vitro and in vivo; however, the sites that are phosphorylated are unknown and the functional significance of phosphorylation has not been well characterized (12). Coronin 1B (also known as CoroninSE or Coronin-2) was discovered as a carbachol-stimulated phosphoprotein in rabbit parietal cells (13). Carbachol acts via a protein kinase C (PKC) pathway in

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this cell type to regulate secretion. Recently, this gene was identified in a microarray-based screen for genes up-regulated in a neural regeneration model (14). Overexpression of Coronin 1B induced neurite extension in several neuronal cell lines, whereas reduced expression via RNA interference caused a reduction in outgrowth. Two-dimensional gel electrophoresis studies suggest that Coronin 1C (Coronin-3) is also phosphorylated in vivo (15).

The potential PKC regulation of mammalian Coronins is intriguing because this family of Ser/Thr kinases regulates migration in many cell types. Activation of classic and novel isoforms of PKC by the phorbol ester phorbo12-myristate 13-acetate (PMA) in some cell types increases cell migration, whereas in other cell types, it is inhibitory (16). These opposing effects are likely because of the complement of PKC isoforms and/or substrate proteins expressed in a given cell type (17). PMA is thought to exert its effects on migration by changing adhesiveness and/or actin architecture at the leading edge (18, 19). PMA also regulates other actin-based motility events such as neurite outgrowth and endosome rocking (20, 21). Despite the clear importance of PKC in cell motility, a relatively small number of PKC substrate proteins have been thoroughly characterized in terms of the residues phosphorylated and functional significance of their phosphorylation in the control of motility. In this work, we have identified serine 2 as the major phosphorylation site on Coronin 1B, and demonstrated that phosphorylation regulates the interaction between Coronin 1B and the Arp2/3 complex. Furthermore, we show that Ser-2 phosphorylation of Coronin 1B regulates PMA-induced ruffling and cell migration.

**EXPERIMENTAL PROCEDURES**

**Materials**—Commercial antibodies were obtained from Cell Signaling Technologies (pSerPKC), Upstate Biotechnology (p84-Arc), Roche (GFP), and Sigma (9E10/Myc). Alexa Fluor 568 phalloidin was from Molecular Probes. Cell lines were from ATCC. PKC inhibitors Ro32-0432 and Go6976 were from Calbiochem. Protease and phosphatase inhibitors (phenylmethylsulfonyl fluoride, 1,10-phenanthroline, aprolin, leupeptin, sodium fluoride, and sodium orthovanadate) were from Sigma. All other materials were from Fisher Scientific unless otherwise indicated.

**TaqMan Quantitative Real-time PCR**—Total RNA was isolated from murine tissues using a Qiagen RNAeasy kit according to the manufacturer’s protocol. Reverse transcription into cDNA was done in a 20-μl volume using oligo(dT)16-Primer, 10 μm dNTP mixture, and ImProm-II Reverse Transcriptase (Promega) according to the manufacturer’s instructions. PCR were carried out in a 20-μl volume consisting of 10 μl of Universal Master Mix No AmpErase UNG (Applied Biosystems), 0.25 μl of fluorescent probe (Mm00486998_m1[11A], Mm00485552_g1[1B], Mm00485558_m1[11C], Mm00619135_m1[12A], Mm0055812_m1[12B], Mm00504152_m1[12POD], Applied Biosystems), and 8 μl of diluted cDNA (~80 ng of total RNA/reaction). The PCRs were carried out under standard conditions in a 20-μl volume in duplicate for each sample in the ABI Prism 7700 Sequence Detection System (Applied Biosystems). The number of PCR cycles needed to reach the fluorescence threshold was determined in duplicate for each cDNA, averaged, and then normalized against a reference gene, 18 S (4335760–0311007 Applied Biosystems), to yield the cycle number at which fluorescence was reached (CT). For the absolute copy number of Coronin 1A, 1B, 1C, 2A, 2B, and POD, 4-fold dilutions of the respective mouse full-length cDNAs were used to generate a standard curve (22). All tissue samples tested were within the linear range (19–38 cycles) of the assay.

**Molecular Cloning**—PCR and subcloning were performed using standard methods. Detailed methods and primer sequences are available upon request. WTB1-EGFP and WTB2-EGFP containing the complete coding sequence of human Coronin 1B and cloning it into a pMSCV-based retroviral vector (23). S2A-EGFP and S2D-EGFP were generated from WTB1-EGFP by site-directed mutagenesis using mutation-encoding primers. Coronin 1B-Myc was generated by subcloning the Coronin 1B coding sequence into a pMSCV vector containing a C-terminal Myc tag.

**Cell Culture, Transient Transfection, and Retroviral Transduction—** HEK293 and Swiss 3T3 cells were cultured in Dulbecco’s modified Eagle’s medium-H supplemented with 10% fetal bovine serum (HyClone), 100 units/ml penicillin, 100 μg/ml streptomycin, and 292 μg/ml glutamine. Rat2 and NIH3T3 cells were cultured in the same media supplemented with 5% fetal BSA serum or 10% calf serum (HyClone), 100 units/ml penicillin, 100 μg/ml streptomycin, and 292 μg/ml glutamine. HEK293 cells were transiently transfected with plasmids using FuGENE 6 transfection reagent (Roche) according to the manufacturer’s protocol. Retroviral packaging, infections, and fluorescence-activated cell sorting were performed as described previously (23).

**Immunoprecipitation**—Cells were washed twice with phosphate-buffered saline and lysed with a KCI buffer (20 mM HEPES, pH 7.0, 100 mM KCl, 0.5 mM EDTA, 0.1% Triton X-100, 0.1% sodium fluoride, 10 μg/ml 1,10-phenanthroline, 10 μg/ml aprolin, 10 μg/ml leupeptin, 10 mM sodium fluoride, and 2 mM sodium orthovanadate). Lysates were cleared at 13,000 × g for 5 min and incubated with 0.5 μg of primary antibody for 1 h at 4 °C, followed by the addition of 20 μl of 50% slurry of ImmunoPure-immobilized protein A/G beads (Pierce) and further incubation at 4 °C for an additional hour. Beads were pre-blocked with 1 mg/ml bovine serum albumin and washed extensively with the KCI buffer prior to usage. The immune complexes were collected, washed with the KCl buffer three times, separated by SDS-PAGE, and transferred to a polyvinylidene difluoride membrane (Bio-Rad) for Western blotting.

**Recombinant Protein Production—**His-tagged Coronin 1B (His1B) was generated by PCR cloning the coding sequence of human Coronin 1B into pQE-80L (Qiagen). GST-tagged full-length (GST1B) or the C-terminal fragment of Coronin 1B (GST1BΔC) were cloned into pGEX-6P1 (Amersham Biosciences). For protein production, cells were grown to mid-log and expression was induced with 0.5 mM isopropyl-1-thio-β-D-galactopyranoside for 6 h at room temperature. Bacteria was harvested and sonicated in lysis buffer (50 mM HEPES, pH 7.4, 0.5% Triton X-100, 0.5 mM NaCl, 5 mM 2-mercaptoethanol, 1 mM phenylmethylsulfonyl fluoride, 10 μg/ml 1,10-phenanthroline, 10 μg/ml aprolin, 10 μg/ml leupeptin, and 10 mM imidazole). His1B was purified by affinity chromatography using nickel-chelated agarose column (Pierce). Lysate, cleared at 20,000 × g for 20 min, was applied to a nickel-chelated agarose column, and washed using the lysis buffer containing 40 mM imidazole. His1B was eluted with 500 mM imidazole in 50 mM HEPES, pH 7.4, 0.5 mM NaCl. GST-tagged proteins were purified by affinity chromatography using immobilized glutathione beads (Pierce) according to the manufacturer’s protocol, and the GST tag was removed by PreScission protease (Amersham Biosciences). The recombinant region of mouse Coronin 1B (394–484 amino acids) was PCR cloned into pMALT2 vector (New England Biolabs). Maltose-binding protein was purified by affinity chromatography using amylose resin (New England Biolabs) according to the manufacturer’s protocol.

**In Vitro Phosphorylation**—2 μg of recombinant protein was incubated with ~0.5 μg of PKCs (Panvera or Sigma) in 100 μl of kinase reaction buffer (7.2 mM HEPES, pH 7.4, 3.6 mM MgCl2, 100 μg/ml aprotinin, 1 mM benzamidine, 1 mM leupeptin, 0.1% Triton X-100, 0.63 mM CaCl2, 0.1 μM ATP, 0.1 μM [γ-32P]ATP, 120 μM phosphatidyserine) containing 10 μCi of (γ32P]ATP for 30 min at 30 °C (24). Reactions were terminated by adding an equal volume of 2× sample buffer. The samples were subjected to SDS-PAGE and transferred to a nitrocellulose membrane for autoradiography and peptide mapping.

**In Vivo Phosphorylation**—Cells were serum starved for 12 h before metabolic labeling, and then incubated with [32P]orthophosphoric acid (Amersham Biosciences) in phosphate-free minimum essential medium (Sigma). After 8 h, cells were treated with or without 100 nM PMA for 30 min and lysed with RIPA buffer (50 mM Tris-HCl, pH 7.3, 150 mM NaCl, 1% IPEGA, 0.5% deoxycholate, 1 mM EDTA, 1 mM phenylmethylsulfonyl fluoride, 10 μg/ml 1,10-phenanthroline, 10 μg/ml aprolin, 10 μg/ml leupeptin, 1 mM α-naphthyl acid phosphate, 10 mM sodium orthovanadate, 1% Triton X-100, 10% glycerol). The supernatants of the activated cell sorting were performed as described previously (23).

**Peptide Mapping**—Peptide mapping was performed as described previously (25, 26). Briefly, appropriate protein bands were cut from the nitrocellulose membrane and digested with 1-isoasleamido-2-phenylthioethyl ketone-treated trypsin (Proteina, Worthington) in 0.05 mM NH4HCO3, pH 7.8, at 37 °C for 24 h. The peptides were washed and dried using a speed vacuum centrifuge, and spotted onto cellulose thin layer chromatography plate (Merck KGaA, Germany) for two-dimensional mapping using pH 8.9 buffer (ammonium carbonate, 10 g/liter) for electrophoresis in the first dimension and phosphochromatography buffer (n-butanol:pyridine:glacial acetic acid:deionized water, 75:50:15:80) for chromatography in the second dimension. After
that the thin layer chromatography plates were dried completely for autoradiography.

Coro1B Antibody Production and Affinity Purification—Rabbits were immunized with a GST fusion protein containing the human Coronin 1B C-terminal region (394–489 amino acids) by Covance. Serum was affinity purified against maltose-binding fusion protein containing the equivalent region of mouse Coronin 1B to ensure cross-reactivity. This myelin basic fusion protein was purified in a buffer containing 10 mM MOPS, pH 7.5, and 60 mM sodium citrate, which was used for direct coupling to UltraLink biosupport media (Pierce) according to the manufacturer’s protocol. The column was washed extensively with 10 mM Tris, pH 7.5, 100 mM glycine, pH 2.5, and 100 mM triethylamine, pH 11.5, with neutralizing washes between. Crude serum was diluted 10-fold in 10 mM Tris, pH 7.5, and passed through the column three times. The column was washed with 10 mM Tris, pH 7.5, and then with 500 mM NaCl, 10 mM Tris, pH 7.5. Antibodies were eluted sequentially with 100 mM glycine, pH 2.5, followed by 100 mM triethylamine, pH 11.5. The antibody-containing fractions were neutralized, combined, and dialyzed against phosphate-buffered saline.

Immunofluorescence Microscopy—For immunofluorescent staining, the cells were fixed, stained, and mounted as described previously (23). Cells were stained in various combinations with Alexa Fluor 568 phalloidin for F-actin (1:400 dilution), p34-Arc (1:400 dilution), or affinity purified Coro1B antibody (1:200 dilution). Images were captured using a Nikon Eclipse E600 with a Nipkow-type spinning disk confocal scan head (Yokogawa CSU-10) attached to an inverted microscope (model IX-81, Olympus) equipped with a ×60/1.45 NA objective, a CCD camera (model C4742–80-12AG, Hamamatsu), and controlled by AQM Advance 6 software (Kinetic Imaging, Ltd.). The extended depth of focus plugin (bigwww.epfl.ch/demos/edf/) for ImageJ was used to combine multiple images from a z-stack (0.1–0.2 μm interval) into a single in-focus image. Images were combined and annotated in Photoshop for presentation.

PMA-induced Ruffling Assay—To quantify ruffling, Rat2 cells were serum-starved overnight, stimulated with 100 nM PMA, fixed at various times as indicated, and stained with Alexa Fluor 568 phalloidin to visualize membrane ruffles. At least 200 cells were counted for each time point.

Single Cell Tracking—Tracking was performed as described previously (23). To increase data throughput, 10 different fields (containing ~10–20 cells per field) were simultaneously recorded using an automated X-Y stage (Prior). To avoid bias in the analysis, every cell in each movie that met the tracking criteria (did not divide, was completely within the field of view for the entire experiment, and did not touch another cell for more than two frames) was tracked using Tracking Analysis software (Kinetic Imaging, Ltd.) with the point-click mode. Cell speed was calculated using the Tracking Analysis software and directional persistence was calculated using custom Matlab macros (a kind gift of B. Harms, MIT). Data were analyzed using PRISM (GraphPad, Inc.) for statistical analysis.

RESULTS

As a first step in understanding the role of mammalian Coronins in cell motility, we used quantitative real-time PCR to compare expression patterns of the genes across mouse tissues. An advantage of this technique is the ability to quantitatively compare expression between genes, as well as across tissues. Our data indicate that the most widely expressed mammalian Coronin gene is CORONIN 1B, which is expressed at high levels across most tissues (Fig. 1B). A full characterization of mammalian Coronin gene expression using this technique will be presented elsewhere. To study Coronin 1B at the protein level, a polyclonal antibody was raised against the unique region and the coiled-coil region of human Coronin 1B (Fig. 1A). HEK293 cells were transiently transfected with all six Coronin genes tagged with GFP at the C terminus. The different Coronins were enriched by immunoprecipitation (IP) using a GFP antibody and blotted with the affinity purified Coro1B antibody. This antibody specifically recognizes Coronin 1B and does not cross-react with the other related Coronins (Fig. 1C). Interestingly, the endogenous Coronin 1B was co-immunoprecipitated (co-IP) with Coronin 1B-GFP. The endogenous protein was co-immunoprecipitated only from cells transfected with Coronin 1B-GFP and not with any of the other Coronin constructs, suggesting only homo-oligomerization had occurred. To confirm this result, we co-transfected cells with the complete set of Coronin genes tagged with GFP and Coronin 1B-Myc. Immunoprecipitation with a Myc antibody co-precipitated only Coronin 1B-GFP and none of the other Coronins (Fig. 1D).

Together, these experiments strongly suggest that Coronin 1B exclusively homo-oligomerizes and does not form complexes with the other Coronins. Whereas other Coronins have been shown to homo-oligomerize via their coiled-coil motif in vivo (27), these data are the first to address the specificity of oligomerization between members of this protein family.

To examine the localization of Coronin 1B, we used our affinity purified antibody for immunofluorescent staining and compared the localization of the endogenous protein to the GFP-tagged form in Rat2 fibroblasts (Fig. 1E). Coronin 1B is localized to the leading edge in fibroblasts, as well as weakly along actin stress fibers. The localization of the GFP-tagged form is indistinguishable from the endogenous protein. This observation, along with the co-IP experiment described above, suggests that Coronin 1B can tolerate tagging with GFP at the C terminus.

In yeast, Coronin and the Arp2/3 complex biochemically and genetically interact, but this interaction has not been demonstrated in mammalian systems (6). To examine this interaction, we first performed immunofluorescent co-localization between Coronin 1B and p34-Arc (a component of the Arp2/3 complex). These two proteins strongly and completely co-localize at the leading edge of fibroblasts (Fig. 2A). To test for a biochemical interaction between Coronin 1B and the Arp2/3 complex, we performed a reciprocal co-IP between Coronin 1B and the p34-Arc subunit of the Arp2/3 complex. We saw a robust, reciprocal co-IP between the endogenous Coronin 1B and p34-Arc in Rat2 fibroblasts (Fig. 2B). This result was also obtained in HEK293, Swiss 3T3, and NIH3T3 cells, suggesting that this interaction is generally observed in many cell types (data not shown). Control rabbit antibodies did not IP either protein (Fig. 2, B and C). We also detected co-IP between Coronin 1B and the Arp2 and p41Arc subunits of the Arp2/3 complex (data not shown), suggesting that our results are not specific for p34-Arc, but reflect an interaction between Coronin 1B and the Arp2/3 complex. As an additional control, we were also able to show co-IP between GFP-tagged Coronin 1B and p34-Arc using GFP antibodies (Fig. 6D). We could not detect actin in the immunocomplex by blotting, but to exclude the possibility that this interaction is simply bridged through F-actin, we pretreated cells with Latrunculin B to disrupt F-actin prior to making the lysate for IP and saw no change in the interaction (data not shown).

Because Coronins have been shown to be PKC substrates in vitro and in vivo (12, 13), we tested whether phosphorylation modulates the interaction between Coronin 1B and the Arp2/3 complex. With 100 nM PMA treatment, this interaction is strongly decreased relative to untreated Rat2 cells (Fig. 2C). To establish that the PMA stimulation was acting through a kinase pathway, we used an anti-phosphoserine antibody raised against a phosphopeptide corresponding to the PKC consensus site. This antibody (pSerPKC) recognizes phosphorylated Coronin 1B immunoprecipitated by our affinity purified Coro1B antibody (Fig. 2C). Mutant analysis as described below confirmed the specificity of this antibody for phospho-Coronin 1B. Further evidence for the specific involvement of PKC in regulating Coronin 1B-Arp2/3 interactions comes from treatment with two well established PKC inhibitors (28, 29). Pretreatment of cells with a pan-PKC inhibitor (Ro32–0432) blocked

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2 L. Cai, N. Holoweckyj, and J. E. Bear, unpublished observations.
PMA-stimulated phosphorylation of Coronin 1B and restored the co-IP with p34-Arc. Interestingly, G6976 (which inhibits the \( \alpha \) and \( \beta \)1 isoforms of PKC) did not block phosphorylation or restore co-IP of p34-Arc, suggesting that a novel isoform of PKC (\( \delta \), \( \varepsilon \), or \( \theta \)) may be responsible for the PMA-induced phosphorylation of Coronin 1B in vivo. Together, these experiments suggest that the interaction between Coronin 1B and the Arp2/3 complex is regulated through PKC phosphorylation of Coronin 1B.

Because the phosphorylation status of Coronin 1B is critical for interaction with the Arp2/3 complex, we sought to identify the residues that are phosphorylated in vivo. As a first step, we did an in vitro kinase assay using purified PKC and His-tagged Coronin 1B.
Coronin-Arp2/3 Interaction Regulated by PKC

Coronin 1B. Recombinant Coronin 1B is strongly phosphorylated in this reaction (Fig. 3A). Similar results were obtained using GST-tagged Coronin 1B (see below). As a positive control, we used a fragment of vinculin that has been reported to be an in vitro PKC substrate (24). Tryptic peptide mapping shows that 4 to 5 sites on Coronin 1B are phosphorylated in vitro (Fig. 3B). Tryptic peptide mapping of chicken vinculin tail (VT; amino acids 881–1135) served as a positive control. PKCo autophosphorylation is indicated by the arrowhead. In vitro phosphorylated His-Coronin 1B subjected to tryptic peptide mapping. The asterisk indicates origin. Spots a–e showed a consistent pattern across multiple experiments. C, GFP-tagged Coronin 1B was immunoprecipitated from 32P metabolically labeled HEK293 cells treated with and without 100 nM PMA (arrow indicates Coronin 1B-GFP). D, tryptic peptide map of a sample from C.

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Comparison of the in vitro and in vivo tryptic peptide maps indicated that the in vivo site did not appear to match any of the in vitro sites. We considered two possible explanations for this observation: first, that PKC does not phosphorylate the site in vitro that is utilized in vivo, or second, that the spot had shifted because of an extra sequence on the peptide from the His or GST tag. Comparison of the tryptic peptide maps of in vivo phosphorylated Coronin 1B and in vitro phosphorylated His1B or (GST)1B provide strong evidence for the latter possibility. We used pairwise mixing of samples to compare the maps of in vitro and in vivo phosphorylated proteins (Fig. 4, A–D). Both the in vitro samples showed spots b, c, d, and e (Fig. 4C) that were absent in the in vivo labeled sample. Furthermore, we discovered that each sample contained a unique spot (a, a’, and a”). It is important to note that, in the case of (GST)1B, the N-terminal GST tag was removed by cleavage prior to the phosphorylation reaction leaving five residual amino acids. Because each sample has a unique N-terminal tryptic peptide because of fusion of the His or GST tags (Fig. 4E), we postulated that a serine or threonine residue common to all of these peptides, Ser-2 in the native protein, might be the in vivo phosphorylated residue. Further evidence for this hypothesis came from the predicted migration properties of the N-terminal peptides. A theoretical peptide map of these peptides, constructed based on established formulas for the relative migration of tryptic peptides, agreed with our experimental observations (Fig. 4F) (25).

To test the hypothesis that Ser-2 is the major in vivo phosphorylation site of Coronin 1B, this residue was mutated to alanine in our GFP-tagged Coronin 1B construct and transfected into HEK293 cells. Upon PMA stimulation, Coronin 1B was strongly phosphorylated (Fig. 3C). Tryptic peptide mapping indicated that there is a single major site phosphorylated in vivo (Fig. 3D).

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correct molecular weight, whereas the S2A sample had no detectable band at the equivalent area of the gel. These samples were blotted with our Coronin 1B antibody to confirm that both proteins were present (data not shown). The tryptic peptide maps generated from the Coronin 1B-GFP bands exhibited a similar pattern of spots as the recombinant preparation studied previously (spots b–e). However, the wild-type sample did not exhibit peptides a or a′/H11033, but rather contained a peptide corresponding to a′/H11032. This spot was absent from the S2A sample (Fig. 5C). Together, these data indicate that Ser-2 is the major site phosphorylated by PKC on Coronin 1B in vivo.

Our experiments using PMA and PKC inhibitors demonstrated that phosphorylation of Coronin 1B regulates its interaction with the Arp2/3 complex. If Ser-2 is the critical phosphorylation site, then mutations in Ser-2 should modulate the Coronin 1B-Arp2/3 interaction. In addition to the S2A mutation described above, we also made a second mutant, S2D, in which Ser-2 was mutated to aspartic acid to mimic phosphorylated Coronin 1B. These mutants, along with the wild-type version, were transiently expressed in HEK293 cells and immunoprecipitated with GFP antibodies. The IP samples were blotted for p34-Arc to detect associated Arp2/3 complex. The
Coronin 1B S2A mutant showed a stronger interaction with the Arp2/3 complex relative to wild type, whereas the S2D mutant had weaker interaction (Fig. 6A). This trend is consistent with our observation that increasing Coronin 1B phosphorylation by PMA treatment decreases the interaction with Arp2/3. Interestingly, these mutants did not affect oligomerization of the GFP-tagged form with the endogenous Coronin 1B, suggesting that this interaction is not regulated by phosphorylation at Ser-2.

To further characterize these mutant versions of Coronin 1B, we created stable Rat2 cell lines expressing the mutants via retroviral transduction. These lines were sorted for equal levels of GFP expression by a fluorescence-activated cell sorter, and equal expression levels of Coronin 1B-GFP were verified by Western blotting (Fig. 6B). For examining phosphorylation status, we used a PMA-induced ruffle formation assay. Previous studies indicated that PMA stimulation of serum-starved cells induces a transient activation of the Rac GTPase and that this Rac activity is required for PMA-induced ruffling activity (30, 31). In our stable Rat2 lines expressing wild-type or Ser-2-mutated Coronin 1B, we observed striking differences in the ability of PMA to induce ruffles in serum-starved cells (Table I). About 41% of the Rat2 control cells show ruffling at 10 min after stimulation, whereas 60% of cells overexpressing wild-type Coronin 1B show ruffling under the same conditions. Both the Rat2 control cells and wild-type Coronin 1B overexpressers had diminished ruffling by 30 min after stimulation, suggesting that this response adapts over time. Cells expressing the S2A mutant version of Coronin 1B had the highest level of ruffling (71%) at 10 min and had sustained high levels of ruffling at all time points tested. Interestingly, cells expressing the S2D mutant version of Coronin 1B had suppressed ruffling with only about 26% of cells showing ruffling activity at 10 min. The S2D cells did show a peak of ruffling activity at 20 min that returned to control levels by 30 min, suggesting that these cells have diminished ruffling rather than a total loss. Together, these data suggest that overexpression of wild-type or S2A Coronin 1B enhances the early phase of the ruffling response, and that Coronin 1B phosphorylation at Ser-2 may be a major factor in the adaptation of PMA-induced ruffling at later times. Furthermore, these data suggest that, consistent with its localization pattern, Coronin 1B exerts a strong effect on actin-dependent processes at the leading edge.

To further explore Coronin 1B function and the role of Ser-2 phosphorylation in regulating Coronin function, we turned to the single cell tracking assay. In this assay, individual cells undergoing random cell movement are tracked in time-lapse movies and information about speed and directional persistence are extracted. Overexpression of wild-type Coronin 1B caused a slight, but statistically significant increase in average cell speed (analysis of variance, $F = 106.7, p < 0.0001$; Dunnett’s multiple comparison test, $p < 0.05$) (Fig. 7A). Expression of the S2A mutant caused a much stronger increase in cell speed, whereas expression of the S2D mutant suppressed cell speed below that of the parental Rat2 cells (Dunnett’s multiple comparison test, $p < 0.01$). None of the cell lines showed significant differences in directional persistence (data not shown).

To understand the relative contribution of Ser-2 phosphorylation to the overall effect of PMA on cell migration, we first examined the dose relationship of PMA treatment to Coronin 1B phosphorylation by Western blotting. In this experiment, the cells were treated with PMA for 30 min in serum-containing media. For examining phosphorylation status, we used Rat2 cells expressing wild-type Coronin 1B tagged with GFP to
examine the phosphorylation of both the GFP-tagged and endogenous forms. Cells were stimulated with different doses of PMA and the Coronin 1B-GFP immunoprecipitated with GFP antibodies and blotted with Coro1B or p34-Arc antibodies. The arrowhead indicates GFP-tagged Coronin 1B, and the arrow indicates endogenous Coronin 1B. B, lysates from Rat2 stable cell lines expressing GFP-tagged wild-type, S2A, or S2D mutant Coronin 1B blotted as indicated. C, GFP IPs of samples in B blotted as indicated. Note the lack of cross-reactivity of pSerPKC antibody with either Ser-2 mutant. D, GFP-tagged Ser-2 mutants have the same localization pattern as the wild-type Coronin 1B in Rat2 cells. Cells were stained with antibodies to p34-Arc to label Arp2/3 and phalloidin to label F-actin.

TABLE I

|               | Untreated | 5 min    | 10 min   | 20 min   | 30 min   |
|---------------|-----------|----------|----------|----------|----------|
|               |           |          |          |          |          |
| Rat2          | 13.4% (46/343) | 38.4% (159/414) | 41.4% (165/399) | 52.2% (224/429) | 38.8% (150/387) |
| WT1B          | 30.1% (64/213) | 55.4% (185/334) | 60.3% (188/312) | 64.2% (280/436) | 47.0% (236/502) |
| S2A           | 17.8% (40/225) | 63.3% (209/330) | 71.3% (268/376) | 76.1% (331/435) | 76.1% (299/393) |
| S2D           | 25.4% (63/248) | 13.2% (39/287) | 26.8% (110/410) | 45.0% (172/382) | 32.8% (168/513) |

FIG. 6. Ser-2 mutants affect the interaction between Coronin 1B and the Arp2/3 complex, but not subcellular localization. A, transient transfection of Ser-2 mutants into HEK293 cells. Samples were immunoprecipitated with GFP antibody and blotted with Coro1B or p34-Arc antibodies. The arrowhead indicates GFP-tagged Coronin 1B, and the arrow indicates endogenous Coronin 1B. B, lysates from Rat2 stable cell lines expressing GFP-tagged wild-type, S2A, or S2D mutant Coronin 1B blotted as indicated. C, GFP IPs of samples in B blotted as indicated. Note the lack of cross-reactivity of pSerPKC antibody with either Ser-2 mutant. D, GFP-tagged Ser-2 mutants have the same localization pattern as the wild-type Coronin 1B in Rat2 cells. Cells were stained with antibodies to p34-Arc to label Arp2/3 and phalloidin to label F-actin.

PMA-induced ruffling response

Data presented are the combined results from two independent experiments. Cells were counted in 6 fields of view with ruffling cells/total, cells counted are reported in parentheses. Chi-square values for each time point range from 38.5 to 82.9, indicating that each time point contains statically significant deviation from expected Rat2 control proportion (p < 0.000001).
Western blot analysis where the sharpest increase in phosphorylation occurred between these two doses. Cells expressing the S2A mutant showed a dose-dependent decrease in cell speed to a point, but the decline in cell speed leveled off by 5 nM and showed no further decrease after this point. This suggests that a mutant version of Coronin 1B that cannot be phosphorylated confers partial resistance to the inhibitory effects of PMA on cell speed. Conversely, the S2D mutant rendered the cells hypersensitive to the inhibitory effects of PMA with the majority of the inhibitory effect occurring by the 5 nM dose. Together, these data indicate that Coronin 1B phosphorylation at Ser-2 plays a significant role in mediating the effects of PMA on cell migration speed in fibroblasts.

DISCUSSION

Coronins are a highly conserved family of proteins that regulate actin-dependent processes such as migration and endocytosis; however, little is known about the function or regulation of these molecules in mammalian systems. In this work, we show that Coronin 1B interacts in vivo with the Arp2/3 complex and that this interaction is inhibited by PKC phosphorylation. We have identified the single major in vivo phosphorylation site, Ser-2, and demonstrated that mutations in this site cause striking effects on fibroblast motility. In addition, we show that Coronin 1B phosphorylation mediates a significant fraction of PMA-induced changes in fibroblast motility.

Mammalian genomes contain at least six Coronin genes that can be broken down into three subclasses based on sequence homology, localization, and function. Coronins 1A, 1B, and 1C form a distinct group and are most similar to the Coronin genes in single-cell eukaryotes. The Coronin 1B and 1C genes are ubiquitously expressed, but Coronin 1B is expressed at 2–10-fold higher levels at the mRNA level depending on the tissue. This suggests that Coronin 1B may be the predominant Coronin of this subclass in many cell types and tissues.

Because Coronin 1B is expressed at such high levels across tissues, we examined a number of molecular interactions that have been seen with other Coronins for this protein. First, our data show that Coronin 1B oligomerizes with itself and that this interaction is exclusively homo-oligomeric in nature. It will be interesting to see if all mammalian Coronins display this degree of specificity in their oligomerization. Second, like all other Coronins tested from any species, recombinant Coronin 1B co-sediments with F-actin in vitro. Third, our reciprocal co-IP and co-localization data show that Coronin 1B interacts in vivo with the Arp2/3 complex in all cell types tested. Furthermore, we see similar interactions between Coronin 1C and Arp2/3. Whereas Coronin 1A was identified as a major co-purifying protein in preparations of the Arp2/3 complex from neutrophils, the data reported in this work are the first to directly confirm that the Coronin-Arp2/3 interaction, first dem-

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3 N. Holoweckyj and J. E. Bear, unpublished observation.
Coronin-Arp2/3 Interaction Regulated by PKC

Our data are the first direct demonstration that phosphorylation regulates Coronin function. Other groups have reported PKC phosphorylation of Coronins 1A and 1B and, for Coronin 1A, there is an increased localization of the protein on phagosomes in HL-60 cells treated with high doses of PKC inhibitors, but none of these studies directly show regulation by phosphorylation (12, 13). Our studies reveal that the in vivo interaction between Coronin 1B and Arp2/3 is strongly inhibited by PKC phosphorylation. We mapped the major in vivo PKC phosphorylation site on Coronin 1B to Ser-2 using tryptic peptide analysis and confirmed this result using site-directed mutagenesis. Expressing Ser-2 mutants has striking effects on fibroblast motility. The S2A mutant has stronger interaction with Arp2/3, and increases PMA-induced ruffling and cell speed relative to the wild-type Coronin 1B, suggesting that this mutant is constitutively active. Conversely, the S2D mutant has reduced interaction with Arp2/3, inhibits ruffling, and suppresses cell speed below that of the parental Rat2 cells, suggesting that this mutation creates a dominant negative Coronin.

Although our data shows regulation of Coronin function by PKC, several outstanding issues remain to be resolved. The precise isoform or isoforms of PKC that phosphorylate Coronin 1B in vivo remain to be delineated. Furthermore, the upstream signaling pathways that ultimately regulate Coronin 1B phosphorylation also remain unclear. In rabbit parietal cells, carbachol (a muscarinic agonist) stimulates Coronin 1B phosphorylation (13), but we were unable to stimulate Coronin 1B phosphorylation in HEK293 cells with this agonist (data not shown). However, this may simply reflect poor/absent expression of the necessary receptor or G-protein in this cell type. A second unresolved issue concerning the PKC regulation of Coronin 1B function is the nature of the dominant negative effect observed with the S2D mutant. At least two possible explanations could account for this effect. First, this pseudo-phosphorylated version may be non-functional, but still able to bind to and sequester key Coronin 1B binding partners. Second, phosphorylated Coronin 1B may have a distinct function from the dephospho-form that does not involve Arp2/3 and is inhibitory for ruffling and migration. More experiments will be needed to resolve these issues.

Whereas the mechanistic details of mammalian Coronin function remain unclear, it is worth considering data on Coronin function from yeast in developing a working hypothesis about Coronin 1B function in mammalian cell motility. In yeast, Coronin binds directly to the Arp2/3 complex in vitro and, more specifically, to the p35-Arc subunit in two-hybrid assays (6). When added to in vitro polymerization assays, Coronin inhibits the Arp2/3-mediated nucleation of actin filaments. Recent structural data indicate that the addition of Coronin to Arp2/3 preparations (visualized by EM and single particle reconstruction) causes Arp2/3 to adopt an open or inactive conformation (33). Together, these data led the Goode laboratory (33) to propose that yeast Coronin is a direct inhibitor of the Arp2/3 complex. Our data indicate that overexpression of the mutant form of Coronin 1B (S2A) that interacts more robustly with Arp2/3 drives the actin-dependent processes of membrane ruffling and cell migration.

Superficially, these datasets seem inconsistent, but there are a number of possible models in which both could be correct or at least consistent within a particular cellular context. One possibility is that Coronin inhibits Arp2/3 activity in both yeast and mammalian cells, but that in fibroblasts, this results (indirectly) in the disassembly of actin networks through other mechanisms such as ADF/cofilin. This would result in accelerated actin network turnover, and that could lead to enhanced cell motility. Such a mechanism would be consistent with the genetic synergy between mutations in the Coronin and Cofilin genes in yeast (5). Another possibility is that Coronin recruits inactive Arp2/3 to the sides of actin filaments at specific locations within cells where it can be acted upon by activators such as SCAR/WASP proteins as previously suggested (6). Phosphorylation of Coronin (and subsequent loss of interaction with Arp2/3) could act as a switch to expose “activable” Arp2/3 that had been previously docked on filaments at specific locations within the cell. Neither of these models, however, completely explains why Coronin is a biochemical inhibitor of Arp2/3, but a cellular “activator” of Arp2/3-dependent processes such as membrane ruffling and cell migration in fibroblasts. Alternatively, mammalian Coronins and yeast Coronin may both interact with Arp2/3, but have evolved divergent mechanisms for regulating the complex. Experiments are underway to rigorously test these and other possible models.

One of our most striking findings is that Coronin 1B phosphorylation mediates a significant fraction of effects of PMA on fibroblast ruffling and migration. This is a somewhat surprising result because PMA activates all of the conventional and novel isoforms of PKC, and likely stimulates the phosphorylation of a large number of protein substrates. When considering our results with PMA, it is important to take into account the precise conditions of treatment and the different phenomenon being observed. The PMA-induced ruffling is highly dependent on serum starvation and may represent a different effect than the 2–3-h treatments in the presence of serum that we used for the cell tracking studies. Our ruffling data are consistent with Coronin 1B playing at least two distinct roles in this process. First, overexpression of wild-type (or S2A) Coronin 1B enhances the PMA-induced ruffling response that is initiated by an unknown PMA target and presumably involves a Rac-dependent pathway. Second, subsequent phosphorylation of Coronin 1B appears to be critical for an adaptation of the PMA-induced ruffling response as demonstrated by the sustained ruffling observed in the cells expressing the S2A mutant. In the cell tracking experiments, PMA was added 30 min before the start of the movies in the presence of serum. This treatment did not induce continuous ruffling, but rather seemed to cause a dose-dependent reduction of the polarized morphology normally seen in migrating fibroblasts. The reduction in cell speed likely arises from the failure to maintain the polarized morphology necessary for effective migration. Expression of the S2A mutant version of Coronin 1B partially protected the cells from this loss of polarity induced by PMA (data not shown). This suggests that, in this assay, Coronin 1B phosphorylation may have an important role in regulating cell polarity. Future experiments will explicitly test this hypothesis.

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