Identification of RanBP 9/10 as Interacting Partners for Protein Kinase C (PKC) γ/δ and the D1 Dopamine Receptor: Regulation of PKC-Mediated Receptor Phosphorylation

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ABSTRACT

We reported previously that ethanol treatment regulates D1 receptor phosphorylation and signaling in a protein kinase C (PKC) δ- and PKC-γ-dependent fashion by a mechanism that may involve PKC δ isozyme-specific interacting proteins. Using a PKC isozyme-specific coimmunoprecipitation approach coupled to mass spectrometry, we report the identification of RanBP9 and RanBP10 as novel interacting proteins for both PKC δ and PKCγ. Both RanBP9 and RanBP10 were found to specifically coimmunoprecipitate with both PKC δ and PKCγ; however, this association did not seem to mediate the ethanol regulation of the PKCs. It is noteworthy that the D1 receptor was also found to specifically coimmunoprecipitate with RanBP9/10 from human embryonic kidney (HEK) 293T cells and with endogenous RanBP9 from rat kidney. RanBP9 and RanBP10 were also found to colocalize at the cellular level with the D1 receptor in both kidney and brain tissue. Although overexpression of RanBP9 or RanBP10 in HEK293T cells did not seem to alter the kinase activities of either PKCδ or PKCγ, both RanBP proteins regulated D1 receptor phosphorylation, signaling, and, in the case of RanBP9, expression. Specifically, overexpression of either RanBP9 or RanBP10 enhanced basal D1 receptor phosphorylation, which was associated with attenuation of D1 receptor-stimulated cAMP accumulation. Moreover, treatment of cells with select PKC inhibitors blocked the RanBP9/10-dependent increase in basal receptor phosphorylation, suggesting that phosphorylation of the receptor by PKC is regulated by RanBP9/10. These data support the idea that RanBP9 and RanBP10 may function as signaling integrators and dictate the efficient regulation of D1 receptor signaling by PKCδ and PKCγ.

Dopamine (DA) is a key signaling molecule in the brain and periphery. The actions of this neurotransmitter are mediated by dopamine receptors, which are seven transmembrane-spanning proteins belonging to the large family of G-protein-coupled receptors. Dopamine receptors are divided into two major subfamilies, referred to as D1-like and D2-like, on the basis of their structure, pharmacology, and function (Missale et al., 1998). The D1-like receptors consist of the D1 and D5 subtypes, which couple to Gs/olf proteins to activate adenyl cyclase and promote the accumulation of intracel-
lular cAMP. In contrast, the D2-like receptors, which consist of the D2, D3, and D4 subtypes, couple to Gαi proteins, which tend to inhibit adenyl cyclase and decrease intracellular cAMP levels.

The D1 receptor is abundantly expressed in the forebrain and it is not surprising that aberrant D1 receptor signaling has been linked to various neuropsychiatric disorders such as substance abuse, schizophrenia, and Parkinson’s disease. For example, ethanol (EtOH) consumption is reduced in genetically modified mice that lack the D1 receptor, or wild-type mice that are administered D1 selective antagonists 7-chloro-8-hydroxy-3-methyl-1-phenyl-2,3,4,5-tetrahydro-1H-3-benzazepine (SCH23390) (El-Ghundi et al., 1998; Price and Middaugh, 2004). Conversely, treatment with D1-selective agonists facilitates EtOH-related behaviors (D’Souza et al., 2003). In addition, D1-selective agonists have been shown to ameliorate the cognitive deficits associated with schizophrenia and improve parkinsonian-like symptoms in animal models (Keibabian et al., 1992; Schneider et al., 1994; McLean et al., 2009). Understanding how D1 receptors are regulated may be useful for future therapeutic interventions.

Receptor phosphorylation is an important post-translational modification that regulates D1 receptor signaling. In particular, receptor phosphorylation generally contributes to desensitization, a process that renders the receptor less sensitive to subsequent agonist stimulation (Kohout and Lefkowitz, 2003). To date, three classes of protein kinases have been reported to phosphorylate the D1 receptor. These include the G protein-coupled receptor kinases (GRKs), AMP-dependent protein kinase (PKA), and protein kinase C (PKC). GRKs generally phosphorylate the D1 receptor under agonist-activated conditions, which results in receptor desensitization (Tiberi et al., 1996; Gardner et al., 2001; Gainetdinov et al., 2004; Rankin et al., 2006). Likewise, PKA also regulates D1 receptor signaling by modulating the rate of agonist-induced receptor desensitization and intracellular trafficking (Ding and Sibley, 1999; Mason et al., 2002). In contrast to GRKs and PKA, very little is known about the regulation of the D1 receptor by PKC phosphorylation. We determined recently that PKC constitutively phosphorylates the D1 receptor and that this negatively regulates dopaminergic signaling (Rex et al., 2008). Moreover, we found that EtOH decreases constitutive PKC phosphorylation of the D1 receptor with a concomitant potentiation of dopaminergic signaling (Rex et al., 2008). It is noteworthy that EtOH was found to directly inhibit the enzymatic activities of PKCγ and PKCδ, but only when they were isolated from the plasma membrane fraction, an effect that was not observed for other PKC isoforms that were tested (Rex et al., 2008). The molecular mechanisms underlying the EtOH-mediated inhibition of membrane-associated PKCγ and PKCδ kinase activities and how they target the D1 receptor are at present unclear. One hypothesis for the membrane-specificity of this effect is that PKC isozyme-specific interacting proteins exist in the plasma membrane and impart EtOH-sensitivity to PKCγ and PKCδ or are themselves the targets of EtOH. It is noteworthy that a growing number of interacting proteins for PKCs have been identified (Staudinger et al., 1997; Rodriguez et al., 1999; Piontek and Brandt, 2003; Zemskov et al., 2003). Likewise, several D1 receptor-interacting proteins have also been reported that include various scaffolding/trafficking proteins (Bermak et al., 2001; Heydorn et al., 2004; Free et al., 2007; Hazelwood et al., 2008).

The aim of the current study was to use a proteomic-based approach to identify and characterize PKCγ/δ isozyme-specific interacting proteins that may be regulated by EtOH and associate with the D1 dopamine receptor. We now report the identification of RanBP9 and RanBP10 as dual-interacting proteins for both PKCγ/δ and the D1 receptor. Although RanBP10 and RanBP9 do not seem to mediate EtOH-dependent inhibition of PKCγ or PKCδ, these interacting proteins were found to modulate D1 receptor activity through increased receptor phosphorylation. Moreover, we found that the RanBP9/10-dependent increase in receptor phosphorylation is blocked by PKC inhibitor treatment, consistent with the idea that these scaffolding proteins may target PKCγ or PKCδ to the D1 receptor.

Materials and Methods

Materials. HEK293-tsa201 (HEK293T) cells were a gift from Dr. V. Ramakrishnan. HA-PKCδ and 3X-FLAG-PKCγ were gifts from Dr. J. M. Mushinski (Mischak et al., 1993) and Dr. A. Newton, respectively. RanBP10-GFP and RanBPM-GFP were gifts from Dr. Guan Wu and Dr. Takeharu Nishimoto, respectively. PKCγ antibody was purchased from GeneTex, Inc. (Irvine, CA). PKCδ antibody was purchased from BD Biosciences (San Jose, CA) and the GFP antibody from Novus Biologicals (Littleton, CO). 3-[3-(dimethylamino)propyl]-5-methoxy-1H-indol-3-yl-[4-(1H-indol-3-yl)-1H-pyrrole-2,5-dione (G66983) and 12-(2-cyanoethyl)-6,7,12,13-tetrahydro-13-methyl-5-oxo-5H-indolodioil(3,4-c)carbazole (G66976) were purchased from Calbiochem (San Diego, CA).

Cell Culture and Transfection. HEK293T cells were cultured in DMEM supplemented with 10% fetal bovine serum, 1 mM sodium pyruvate, 50 μg/ml streptomycin, 50 U/ml gentamicin at 37°C in 5% CO2. HEK293T cells were seeded in 150-mm culture dishes and transfected 24 h later using calcium phosphate precipitation (Clontech, Mountain View, CA).

Commmunoprecipitation and Immunoblot Analysis from HEK293T Cells. To identify candidate PKC-interacting proteins that might be regulated by EtOH, HEK293T cells expressing FLAG-PKCγ, HA-PKCδ, or empty vector were incubated in the presence or absence of EtOH (100 mM/10 min). Cells were harvested in PBS, centrifuged at 250g for 10 min and the pellets homogenized in 1 ml of homogenization buffer (250 mM sucrose, 50 mM Tris-HCl, 10 mM EGTA, 2 mM EDTA, 30 mM NaF, and 20 mM sodium pyrophosphate) supplemented with Mini Complete protease inhibitor cocktail (Roche Applied Science, Indianapolis, IN). The samples were centrifuged at 20,000g for 30 min at 4°C and the supernatant was discarded. The pellet was resuspended and homogenized with 1 ml of solubilization buffer containing 1% Triton X-100 and supplemented with MiniComplete protease inhibitor cocktail. The homogenate was centrifuged for 5 min at 9000g, and the supernatant was used as the “solubilized” membrane fraction. HA-PKCδ or FLAG-PKCγ was immunoprecipitated from the solubilized fraction using either HA-agarose gel or FLAG-M2 gel (Sigma-Aldrich, St. Louis, MO), respectively, and rotated overnight at 4°C. To identify candidate PKC-interacting proteins, isozyme-specific immunoprecipitates were washed three times with PBS and then sent to Applied Biosciences (Hayward, CA) for two-dimensional differential in-gel electrophoresis. In brief, the samples were covalently linked to CyDye Fluors and separated by isoelectric focusing in the first dimension and SDS-PAGE in the second dimension. Image analysis was performed using DeCyder software (GE Healthcare, Chalfont St. Giles, Buckinghamshire, UK), and selected spots.
were chosen for mass spectrometry analysis (matrix-assisted laser desorption ionization/time of flight).

To verify the association of the PKC isoforms or the D1R with RanBP10 and RanBP9, HEK293T cells coexpressing HA-PKCa, FLAG-PKCγ, or FLAG-D-R with vector, RanBP10-GFP or RanBP9-GFP was subjected to commounprecipitation experiments as described above. The resulting immunoprecipitates were washed three times with PBS, resuspended in SDS-PAGE sample buffer, and incubated for 1 h at 37°C. Proteins were separated electrophoretically using 4 to 12% Bis-Tris NuPAGE gels/MOPS buffer (Invitrogen, Carlsbad, CA) and transferred to polyvinylidene fluoride (PVDF) membranes (Invitrogen). Membranes were blocked in SuperBlock (Thermo Fisher Scientific, Waltham, MA) and washed with TBS-T. Membranes were incubated with primary antibodies. The blots were washed three times with 20 mM Tris-HCl, pH 7.5, 150 mM NaCl, and 0.05% Tween 20, incubated with horseradish peroxidase-conjugated secondary antibodies (Santa Cruz Biotechnology, Santa Cruz, CA), and visualized using SuperSignal Chemiluminescence Kit (Thermo Fisher Scientific). The optical density of the immunoblots was quantified by ImageJ software (http://rsweb.nih.gov/ij/).

In Vitro PKC Assays. HEK293T cells coexpressing FLAG-PKCγ or HA-PKCa8 with either vector, RanBP10-GFP, or RanBP9-GFP were commounprecipitated from the solubilized membrane fraction as described above. Immunoprecipitates were centrifuged at 9000 × g for 1 min at 4°C, and the pellets were washed three times with PBS. Each pellet was resuspended with 70 μl of PBS. Kinase assays were performed using a PKC Assay Kit (Millipore, Billerica, MA) as described previously (Rex et al., 2008). In brief, the kinase activity in 100 μl of each sample was assayed by measuring the transfer of 32P from [γ-32P]ATP to a specific substrate peptide (Millipore). The remaining cell lysates were retained and used to determine PKC expression for each transfection. PKC activity was measured in an assay consisting of assay dilution buffer II (end concentration, 3.33 mM MOPS, pH 7.2, 4.2 mM β-glycerol phosphate, 0.17 mM sodium orthovanadate, 0.17 mM dithiothreitol, and 0.12 mM CaCl2), 83 μM PKC substrate peptide, 0.3 μM PKA inhibitor peptide, 3.3 μM Ca2+/calmodulin-dependent protein kinase inhibitor, PKC lipid activators [80 ng/ml phosphatidyl serine (PS) and 8 ng/ml diacylglycerol (DAG)]. The addition of EtOH was made from a concentrated stock solution. Maximal PKC activity was achieved in the presence of PS and DAG. Basal activity was measured in the presence of 0.5 mM EGTA instead of PS and DAG. Nonspecific activity was determined in the absence of substrate peptide. Each condition was performed in duplicate. Kinase reactions were initiated by the addition of 10 μCi [γ-32P]ATP in Mg2+/ATP cocktail and incubated for 10 min at 30°C. The reactions were terminated by transferring 25 μl of the mixture to P81 filter papers followed by washing three times with 0.75% phosphoric acid and once with acetone. The amount of phosphorylated peptide was determined by scintillation counting and expressed as the percentage of control. To compare PKC activity between the three groups (PKCγ + vector, PKCγ + RanBP10, and PKCγ + RanBP9), the remaining cell lysates were subjected to Western blot and densitometry analysis (ImageJ) to determine the relative amounts of PKC protein. To account for any changes in PKC expression between each cotransfection, the kinase activity was divided by the relative PKC expression (arbitrary units).

cAMP Accumulation Assays. Transfected HEK293T cells were seeded into 24-well plates coated with poly(b-lysine). Duplicate wells were exposed to dopamine dilutions that were prepared in 20 mM HEPES-buffered DMEM supplemented with 200 μM sodium metabisulfite and 30 μM 4-((3-butoxy-4-methoxyphenyl)-methyl)-2-imidazolidinone (Ro 20-1724) (a phosphodiesterase inhibitor) (Sigma-Aldrich). Basal activity was determined in the absence of dopamine. The plates were incubated at 37°C for 20 min. The reaction was terminated by warming the plates to the reaction temperature in 1 N hydrochloric acid to each well for 30 min on ice. Each reaction was neutralized by the addition of 15% KHCO3. cAMP accumulation was measured using the method described by Watts and Neve (1996) based on the competitive binding of cAMP to PKA.

Radioligand Binding Assays. D1 receptor-transfected HEK293T cells were harvested in Ca2+/Mg2+-free Earle’s balanced salt solution supplemented with 5 mM EDTA and centrifuged at 200g for 10 min. Cells were lysed in a Dounce homogenizer in 5 mM Tris-HCl, pH 7.4 (at 4°C) and 5 mM MgCl2. The lysate was centrifuged at 20,000g for 30 min, and the pellet was resuspended in 50 mM Tris-HCl (crude membrane fraction). A portion of the membrane suspension was quantified using the BCA protein assay kit. The membrane fraction (100 μl) was added to tubes in triplicate containing [3H]SCH23390 (PerkinElmer Life and Analytical Sciences, Waltham, MA) at a range of concentrations. Nonspecific binding was determined in the presence of (+)butaclamol (3 μM). Assay tubes were incubated at room temperature for 1.5 h and then terminated by rapid filtration through GF/C filters pretreated with 0.6% polyethyleneimine. Bound radioactivity was quantified by liquid scintillation counting.

In Situ Phosphorylation Assays. These assays were performed as described previously (Rankin et al., 2006). In brief, transfected HEK293T cells were seeded into six-well plates coated with poly(lysine). A portion of the transfection was retained in a 100-mm dish for radioligand binding assay to quantify the expression of D1 receptor expression. Forty-eight hours after transfection, the medium from each well was replaced with phosphate-free medium supplemented with 10% fetal bovine serum (FBS), 50 U/ml penicillin, and 10 μg/ml gentamicin and incubated for 1 h. The medium was replaced with 1 ml of phosphate-free DMEM containing 106 μCi/ml [32P]orthophosphoric acid (PerkinElmer Life and Analytical Sciences) for 60 min. PKC inhibitors (Gö6983 and Gö6976 at 10 μM) were included during this incubation period for select samples. Subsequently, specific wells were challenged with basal medium or media containing 10 μM dopamine for 10 min. The cells were placed on ice, rinsed with ice-cold Earle’s balanced salt solution, and lysed with solubilization buffer supplemented with MiniComplete protease inhibitor cocktail for 1 h at 4°C. Cell lysates were cleared by centrifugation, and protein concentration was quantified using the BCA protein assay kit. D1 receptor expression was determined by radioligand binding assays using cells seeded in the 100-mm dish. Equal amounts of D1 receptor for each condition were incubated with anti-FLAG-M2 agarose gel overnight, washed three times, and resolved using 4 to 12% Bis-Tris NuPage gels using MOPS buffer. Dried gels were subjected to autoradiography.

Immunofluorescence and Confocal Studies of Proximal Tubule Cells in Rat Kidney. Normotensive 12-week-old Wistar-Kyoto rats were anesthetized with pentobarbital (50 mg/kg i.p.). The kidneys were perfused with normal saline until the effluent was clear. Thin sections (3 μm) of formalin-fixed and paraffin-embedded rat kidney were deparaffinized in xylene and then rehydrated with step-down concentrations of ethanol. After antigen retrieval with citrate buffer, the D1 receptor was visualized using a monoclonal mouse anti-D1 receptor antibody (1:100; Novus) followed by Alexa Fluor 568-donkey anti-mouse IgG antibody (red; Invitrogen). RanBP9 was visualized using a polyclonal goat anti-RanBP9 antibody (1:100; Santa Cruz Biotechnology) followed by Alexa Fluor 488-donkey anti-goat IgG antibody (green; Invitrogen). As a negative control, the primary antibodies were replaced with normal mouse, rabbit, or goat IgG at an appropriate dilution. The slides were mounted in Vectashield Mounting Medium with 4,6-diamidino-2-phenylindole (Vector Laboratories, Burlingame, CA). The immunofluorescent images were acquired using Zeiss 510 confocal laser scanning microscope (Carl Zeiss GmbH, Jena, Germany). Colocalization of D1 receptor and RanBP9 is indicated by the development of a yellow color in the merged images.

D1 Receptor and RanBP9 Coimmunoprecipitation from Kidney Cells. Immortalized renal proximal tubule cells (passage ~20) from adult normotensive adult humans and Wistar-Kyoto rats were used for coimmunoprecipitation (coIP). The renal proximal tubule cells were cultured at 37°C in 95% air/5% CO2 atmosphere in
DMEM/F-12 with transferrin (5 μg/ml), insulin (5 μg/ml), epidermal growth factor (10 ng/ml), dexamethasone (4 μg/ml), and 5% FBS in a 100-mm Petri dish. The cells (80% confluence) were made quiescent by incubation for 2 h in medium without FBS before the treatment with a D1-like receptor agonist, fenoldopam (1 μM, 10 min). The cells were lysed with ice-cold lysis buffer for 1 h and centrifuged at 16,000g for 30 min. Equal amounts of cell lysates (supernatant; 500 μg) were used for coIP. coIP was performed according to the manufacturer’s immunoprecipitation protocol (Santa Cruz Biotechnology). In brief, 2 μg of rabbit D1 receptor antibody (D, R 408), the specificity of which has been established previously (Yu et al., 2006), was incubated with immunoprecipitated matrix at 4°C on a rotator for 1 h, and then the complex was incubated with 50 μg of lysate protein at 4°C overnight. The immunoprecipitate was pelleted and washed four times with phosphate-buffered saline. The pellet was resuspended in sample buffer, boiled, and immunoblotted with the rabbit RanBP9 antibody. To determine the specificity of the bands, normal rabbit IgG was used as negative control, and cell lysate (immunoblot) was used as positive control.

Identification of RanBP9 Expression in D1 Receptor-Expressing Neurons in Mouse Striatum. Immunohistochemical localization of RanBP9 and RanBP10 was performed to determine possible coexpression in neurons in the striatum that express the D1 receptor. Coronal brain sections were processed from mice in which a D1 receptor transgene had been perfusion-fixed with 4% formaldehyde, postfixed overnight, and disrupted, and the PKC constructs were immunoprecipitated (IP) from HEK293T cells, whereas untransfected cells served as controls. The cells were identified based on comparing peptide sequences determined by scanning and analyzing the bands using ImageJ software.

Results
Identification of PKCγ and PKCδ-Interacting Proteins. We reported previously that EtOH treatment decreased the kinase activities of membrane-associated PKCγ and PKCδ under lipid-activated conditions (Rex et al., 2008). In contrast, cytosolic PKCγ and PKCδ activities were unaffected by EtOH (Rex et al., 2008). One hypothesis for the membrane-specificity of this effect is that PKC isozyme-specific interacting proteins exist in the plasma membrane and impart EtOH-sensitivity to PKCγ and PKCδ or are the targets of EtOH themselves. To address this hypothesis, we used a PKC isozyme-specific coimmunoprecipitation approach coupled to mass spectrometry sequence analysis to identify novel PKCγ- and PKCδ-interacting proteins, including those that might be regulated by EtOH.

The proteomics approach that we used is illustrated in Fig. 1. FLAG-tagged PKCγ was immunoprecipitated from solubilized membrane fractions prepared from HEK293T cells that were either untreated or treated with EtOH for 15 min. As a control, untransfected HEK293T cells were used for immunoprecipitation with anti-FLAG agarose beads. Likewise, HA-tagged PKCδ was immunoprecipitated from transfected cells using HA-agarose beads under identical conditions as described above. The precipitates for each condition (control, PKCγ, and PKCδ) were separately covalently labeled with fluorochromes (Cy2, Cy3, and Cy5) and subjected to twodimensional differential in-gel electrophoresis. Proteins were separated by isoelectric focusing in the first dimension and SDS-PAGE in the second dimension. This method allows three samples to be run on a single gel and comparison of each sample based on the wavelength of the specific fluorochrome.

Figure 2, A and D, shows a comparison of immunoprecipitated proteins from control (untransfected) versus PKC-transfected cells. In this comparison, any unique proteins that are identified in the PKC-transfected samples are tentatively identified as specific interacting proteins for those PKCs. From the PKCγ screen, we identified seven protein spots that seemed to be specific for PKCγ (red spots) (Fig. 2A). We were also interested in identifying any immunoprecipitated proteins in the PKC-transfected samples that increased or decreased in intensity upon treatment of the cells with EtOH. Comparing PKCγ versus PKCγ + EtOH-treated samples, a single red protein spot was present only in the EtOH-treated sample (Fig. 2B). Furthermore, this protein spot was not present in control (untransfected) cells, suggesting that the association of this protein with PKCγ might be regulated by EtOH (Fig. 2C). We identified five additional protein spots from the PKCδ + EtOH-treated samples that also seemed to be PKCδ-specific, one of which might be up-regulated by EtOH treatment (red spot, Fig. 2E). All of the circled protein spots in Fig. 2 were excised and subjected to mass spectrometry sequence analysis.

The candidate PKC isozyme-specific interacting proteins were identified based on comparing peptide sequences de-
rived from the mass spectrometry analyses with protein sequences submitted to the National Center for Biotechnology Information database. In some instances a single spot on the gel corresponded to multiple parent proteins. For the PKCγ screen, eight protein spots were sequenced that corresponded to 31 different parent proteins; however, in many instances, only a single peptide sequence for that protein was identified. In general, we only considered proteins as bona fide if they were identified through multiple matching peptide fragments. Table 1 shows a list of such proteins obtained from the PKCγ and PKCδ proteomic screens. A variety of different proteins were identified ranging from scaffolding proteins to other protein kinase C isozymes. It is noteworthy that 70-kDa heat shock protein (Hsp-70) was the only candidate PKC-interacting protein that was common to both PKC screens. Heat shock proteins assist in protein folding and assembly and are frequently observed in proteomic screens, usually as nonspecific interactors.

Based on the sequence data for the protein spot excised from the PKCγ + EtOH-treated sample (Fig. 2B), 11 nonoverlapping peptide fragments were identified that correspond to the ephrin receptor EphB1 (Holder and Klein, 1999). Additional peptide fragments identified from this protein spot corresponded to the tripartite motif protein TRIM5 (Johnson and Sawyer, 2009). From the PKCδ screen, the single spot identified in the EtOH-treated samples (Fig. 2E) corresponds to mono-ADP-ribosyltransferase (Okada et al., 2008). Although the potential regulation of these three proteins by EtOH and their interactions with PKCγ and PKCδ should be investigated further, our initial assumption is that they are not responsible for the unique membrane-specific regulation of PKCγ and PKCδ activities observed previously (Rex et al., 2008), because none of them were identified in both PKC screens. Thus, while continuing to pursue the mechanism of EtOH-mediated regulation of PKCγ and PKCδ activities, we chose to initially focus on characterizing other candidate PKC-interacting proteins that might be especially relevant to regulating D1 receptor function. One such protein, RanBP10, and a closely related protein, RanBP9, were chosen for further characterization because RanBP9 has been shown to interact with other GPCRs (see below).

**RanBP10 and RanBP9 Interact with PKCγ and PKCδ.** RanBP10 was among the candidate PKCγ-interacting proteins from our proteomic screen (Fig. 2A and Table 1). It is noteworthy that seven nonoverlapping peptide fragments of RanBP10 were identified by mass spectrometry sequence analysis, all of which span the majority of the protein sequence (Table 2). RanBP10 is a 67-kDa scaffolding protein

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**Table 1**

Candidate PKCγ and PKCδ interacting proteins identified using mass spectrometry analysis

<table>
<thead>
<tr>
<th>Candidate Proteins</th>
<th>Function</th>
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<tbody>
<tr>
<td>RAC/CDC42 exchange factor (GEFT)</td>
<td>Cell signaling</td>
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<tr>
<td>Ran-binding protein 10 (RanBP10)</td>
<td>Scaffold protein</td>
</tr>
<tr>
<td>Hsp-70&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Protein folding/stability</td>
</tr>
<tr>
<td>Glial fibrillary acidic protein</td>
<td>Cell structure/communication</td>
</tr>
<tr>
<td>PKCa</td>
<td>Cell signaling</td>
</tr>
<tr>
<td>Ephrin receptor EphB1&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Cell signaling</td>
</tr>
<tr>
<td>Tripartite motif protein&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Protein localization</td>
</tr>
<tr>
<td>Voltage-gated sodium channel subtype III</td>
<td>Cell signaling</td>
</tr>
<tr>
<td>PKCδ-interacting</td>
<td>Trafficking</td>
</tr>
<tr>
<td>Cytoplasmic dynein</td>
<td>Protein folding/stability</td>
</tr>
<tr>
<td>Hsp-70&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Protein folding/stability</td>
</tr>
<tr>
<td>Hsp-90</td>
<td>Ca&lt;sup&gt;2+&lt;/sup&gt; signaling</td>
</tr>
<tr>
<td>Inositol trisphosphate receptor type 1</td>
<td>Cell structure</td>
</tr>
<tr>
<td>Microtubule-actin crosslinking factor</td>
<td>Protein modification</td>
</tr>
<tr>
<td>Mono-ADP-ribosyltransferase&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Regulation of cellular activities</td>
</tr>
<tr>
<td>Valosin-containing protein</td>
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<sup>a</sup> Common to both PKCγ and PKCδ proteomics screens.

<sup>b</sup> Identified in EtOH-treated samples only.

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Fig. 2. Identification of candidate PKC-interacting proteins. Each CyDYE-labeled PKC sample was simultaneously separated on a 2D-gel and scanned at specific wavelengths to reveal each of the CyDYE signals. Overlay of images reveal protein spots that are presumably specific (or nonspecific) to a given sample. Circled, protein spots selected for identification by mass spectrometry. A to C, PKCγ; D to F, PKCδ.
that is expressed in a variety of tissues including the brain (Wang et al., 2004; Haase et al., 2008; Schulze et al., 2008) and is known to interact with several proteins that include the protein Ran and the receptor protein tyrosine kinase for hepatocyte growth factor, MET (Wang et al., 2004). It is noteworthy that there is a closely related protein, RanBP9, that has high sequence similarity to RanBP10 (Wang et al., 2004) and has been found to associate with several GPCRs, including the μ-opioid receptor and the metabotropic glutamate receptors (mGluRs) mGluR2 and mGluR8, as well as with MET (Wang et al., 2004; Seebahn et al., 2008; Talbot et al., 2009). Figure 3 shows a structural comparison of RanBP9 and RanBP10 highlighting several conserved protein-protein interactions and functional domains such as dual-specific kinase splA and ryanodine receptor, lissencephaly type-1-like homology domain, Ran protein binding domain, and guanine nucleotide exchange factor domain (Schulze et al., 2008). A β-tubulin binding domain has been localized to the N-terminal region of RanBP10 (Schulze et al., 2008). One major difference between RanBP10 and RanBP9 is the presence of a poly proline/glutamine tract at the N terminus of RanBP10 (Fig. 3), the role of which is unclear.

To initially verify the interaction of RanBP10 with PKCγ, we used a communoprecipitation approach as shown in Fig. 4. HEK293T cells were cotransfected with FLAG-PKCγ and either empty vector or RanBP10-GFP. FLAG-PKCγ was immunoprecipitated from solubilized membrane fractions followed by one-dimensional SDS-PAGE electrophoresis and transference to PVDF membranes. The association of RanBP10 with PKCγ was visualized by Western blot analysis. In addition, the effect of EtOH treatment on the association of RanBP10 with PKCγ was examined. For these experiments, cells were pretreated with EtOH for 15 min before the preparation of the membrane fraction and subsequent immunoprecipitation. We confirmed the interaction of RanBP10 with PKCγ using this approach (Fig. 4A). Furthermore, the RanBP10-PKCγ association was not modulated by EtOH treatment. These experiments were also extended to examine the possible association of RanBP10 with PKCδ, even though RanBP10 was not identified in the initial PKCδ proteomic screen. It is noteworthy that RanBP10 was also found to communoprecipitate with PKCδ, and this interaction does not seem to be modulated by EtOH either (Fig. 4B).

Based on the high amino acid sequence similarity of RanBP10 and RanBP9, we wondered whether RanBP9 might also associate with PKCγ or PKCδ. As shown in Supplemental Fig. 1, RanBP9 was also found to communoprecipitate with both PKCγ and PKCδ, but EtOH treatment had no significant effect on this interaction.

**Effect of RanBP10 and RanBP9 on PKCγ and PKCδ Kinase Activities.** Although EtOH treatment did not seem to alter the association of either RanBP10 or RanBP9 with PKCγ or PKCδ (Figs. 4, Supplemental Fig. 1), given that both RanBP proteins interacted with both PKCs, we wondered whether RanBP10 and/or RanBP9 could still somehow be involved in the EtOH-mediated decrease in PKC activity. To directly test this possibility, we used in vitro kinase assays to evaluate PKC activities as we had performed previously (Rex et al., 2008). Our working hypothesis was that if either

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**Table 2**

<table>
<thead>
<tr>
<th>Trypsin-Digested Peptides</th>
<th>Location (RanBP10)</th>
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<tbody>
<tr>
<td>FYAVNQETPLPERSWSFK</td>
<td>42–60</td>
</tr>
<tr>
<td>VNYIGLSQGNLRVHYK</td>
<td>63–78</td>
</tr>
<tr>
<td>ATIP1PAACCTYTFEVK</td>
<td>82–108</td>
</tr>
<tr>
<td>GRDGVGMGIGLSAQVMNRLPGWDK</td>
<td>113–137</td>
</tr>
<tr>
<td>QFVEMVNGTDSEVR</td>
<td>336–349</td>
</tr>
<tr>
<td>RQLCQGNGQATER</td>
<td>499–510</td>
</tr>
<tr>
<td>EPVCAALNISALESQNLPK</td>
<td>566–584</td>
</tr>
</tbody>
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**Fig. 3.** Structural organization of RanBP10 and RanBP9. The RanBP10 and RanBP9 proteins were aligned and compared. Conserved regions include a dual-specific kinase splA and ryanodine receptor, lissencephaly type-1-like homology, Ran protein binding domain, and guanine nucleotide exchange factor domain. RanBP10 contains a poly(proline/glutamine) tract at the amino terminus that is absent in RanBP9. [Adapted from Rex et al., 2008:587–593. Copyright © 2008 The Company of Biologists, Ltd. Used with permission.]

**Fig. 4.** Association of RanBP10 with PKCγ and PKCδ. Communoprecipitation and immunoblot analyses of RanBP10 with PKCγ (A) or PKCδ (B). HEK293T cells were transfected with empty vector, FLAG-PKCγ + RanBP10-GFP, or HA-PKCδ + RanBP10-GFP. PKCγ or PKCδ was immunoprecipitated from solubilized membrane fractions prepared from HEK293T cells that were either untreated or pretreated with ethanol (100 mM/15 min). As a control, the solubilized membrane fraction prepared from cells transfected with empty vector was incubated with either FLAG or HA beads. After SDS-PAGE, the gels were blotted with antisera directed to GFP (top) or to PKCγ or PKCδ. The experiment shown is representative of three independent experiments.
RanBP9 or RanBP10 were involved in imparting EtOH sensitivity to the PKCs, then overexpressing either RanBP protein might enhance the EtOH inhibition. PKC\(\alpha\) or PKC\(\gamma\) were coexpressed with RanBP10, RanBP9, or empty vector in HEK293T cells, followed by immunoprecipitation of the PKCs from a solubilized membrane preparation. The PKC immunoprecipitates were then used directly in the kinase assays. As we determined previously (Rex et al., 2008), the addition of EtOH to the kinase assays decreased the lipid-activated PKC\(\gamma\) and PKC\(\delta\) activities without affecting basal enzyme activities (Fig. 5, A and C). In contrast, we observed that overexpression of either RanBP9 or RanBP10 blocked the EtOH-mediated inhibition of PKC activities without affecting basal or lipid-activated activities (Fig. 5). Taken together, these results suggest that neither RanBP9 nor RanBP10 imparts sensitivity of PKC\(\gamma\) or PKC\(\delta\) to inhibition by ethanol but rather they can inhibit this response, perhaps by blocking the association of the PKCs with another distinct interacting protein.

**RanBP10 and RanBP9 Associate with the D\(_1\) Receptor.** Although RanBP10 and RanBP9 do not seem to mediate the EtOH-induced attenuation of PKC\(\gamma\) and PKC\(\delta\) kinase activities, they could function as important PKC scaffolding proteins. In fact, a role for RanBP10 and RanBP9 as scaffolding proteins has been well described previously (Wang et al., 2004; Haase et al., 2008; Talbot et al., 2009). In particular, both Ran10 and RanBP9 have been reported to associate with receptors expressed at the plasma membrane that include the MET tyrosine kinase receptor and the \(\mu\) opioid receptor, respectively (Wang et al., 2004; Talbot et al., 2009). This was of particular interest because the \(\mu\) opioid receptor is a G protein-coupled receptor, as is the D\(_1\) dopamine receptor. Furthermore, the D\(_1\) receptor is known to be phosphorylated by PKC (Gardner et al., 2001; Rex et al., 2008), which is, at least partially, mediated by PKC\(\delta\) and PKC\(\gamma\) (unpublished observations). We hypothesized that either RanBP10 or RanBP9 might also associate with the D\(_1\) receptor and function as a scaffolding protein, perhaps to position PKC\(\gamma\) or PKC\(\delta\) in close proximity to the receptor. To examine this possibility, we used coimmunoprecipitation analyses to determine whether either RanBP9 or RanBP10 could associate with the D\(_1\) receptor (Fig. 6). HEK293T cells were transfected with a FLAG-D\(_1\)R construct and either RanBP10, RanBP9, or empty vector. A crude membrane preparation was prepared, and the D\(_1\) receptor was immunoprecipitated from the solubilized membrane preparation, separated by one-dimensional SDS-PAGE, and transferred to PVDF membranes. The presence of the D\(_1\) receptor and both RanBP proteins were detected by Western blotting. It is noteworthy that both RanBP10 and RanBP9 were found to specifically coimmunoprecipitate with the D\(_1\) receptor (Fig. 6). These results suggest that RanBP9 and RanBP10 are indeed associated with the D\(_1\) receptor when expressed in HEK293T cells.

**RanBP9 Associates with the D\(_1\) Receptor in Kidney Tissue.** We were interested in examining the association of RanBP9 and RanBP10 with the D\(_1\) receptor in endogenous expressing tissues. Unfortunately, only the RanBP9 antisera proved useful in kidney tissue. To determine whether RanBP9 and the D\(_1\) receptor associate endogenously, we first examined the cellular localization of these proteins in the rat kidney proximal convoluted tubule, in which the D\(_1\) receptor is expressed (Fig. 7). Both RanBP9 and the D\(_1\) receptor were observed to be colocalized in the brush border/apical membrane of the proximal convoluted tubule of the rat kidney (Fig. 7). The D\(_1\) receptor is abundantly expressed in the proximal tubule of humans and rodent kidneys, in which it negatively regulates renal sodium transport at the brush border/apical and basolateral membrane (Zeng et al., 2008). We next attempted to coimmunoprecipitate RanBP9 and the D\(_1\) receptor from renal proximal tubule cells. Figure 8 shows

![Fig. 5. Effect of RanBP9 overexpression on ethanol-depend-ent attenuation of PKC activities. Epitope-tagged PKCs were immunoprecipitated from solubilized membrane fractions prepared from HEK293T cells coexpressing either PKC\(\gamma\) (A and B) or PKC\(\delta\) (C and D) with vector, RanBP10, or RanBP9. The kinase activities of the isozyme-specific immunoprecipitates were directly assessed using an in vitro kinase assay as described under Materials and Methods. For each condition, the kinase activities were measured under basal or lipid-activated (plus phosphatidylserine and diacylglycerol) conditions in the presence or absence of 100 mM ethanol. Data are presented as the mean ± S.E.M. of at least three independent experiments. Results that are significantly different from the control groups are indicated as *, \(p < 0.05\); **, \(p < 0.01\); paired Student’s t test.](https://www.aspetjournals.org/content/L license/agreement)
an experiment in which the D₁ receptor was immunoprecipitated from solubilized renal proximal tubule cells, resolved by one-dimensional SDS-PAGE, and then blotted with antisera directed against RanBP9. Lane 1 shows that a protein corresponding in size to RanBP9 is present in the immunoprecipitates but absent if normal rabbit IgG is used instead of the RanBP9 antisera (Fig. 8, lane 3). This experiment was also performed subsequent to treating the cells with a D₁ agonist; however, this did not alter the association of RanBP9 with the D₁ receptor (Fig. 8, lane 2). Taken together, these experiments show that RanBP9 and the D₁ receptor endogenously associate in the kidney.

RanBP9/10 and the D₁ Receptor Exhibit Cellular Co-localization in Brain Tissue. We were also interested in examining the association of RanBP9/10 and the D₁ receptor in the brain. In this study, immunohistochemical methods demonstrated widespread RanBP9 and RanBP10 immunoreactivity in the forebrain, including the majority of neurons in the cerebral cortex, olfactory bulb and striatum. This pattern of distribution of RanBP9 and RanBP10 immunoreactivity matches that reported for the mRNA for these genes as illustrated in the Allen brain atlas (available at http://www.brain-map.org/). In the striatum, half of the medium spiny neurons express the D₁ receptor, which were visualized directly by their expression of EGFP in the brains of mice used for immunohistochemical localization in this study (Gong et al., 2007). As shown in Fig. 9, immunoreactivity for both RanBP9 (top) and RanBP10 (bottom), was present in striatal neurons. In addition, both antigens were present in D₁-EGFP-positive neurons (yellow arrows) and D₁-EGFP-negative neurons (blue arrows). These data demonstrate that both RanBP9 and RanBP10 colocalize in striatal neurons expressing the D₁ receptor.

RanBP10 and RanBP9 Decrease D₁ Receptor Signaling. To determine whether RanBP9 or RanBP10 association with the D₁ receptor exhibits functional consequences, we examined D₁ receptor-stimulated cAMP accumulation in the absence or presence of RanBP9/RanBP10 overexpression (Fig. 10). It is noteworthy that coexpression of either RanBP10 or RanBP9 significantly decreased maximal DA-stimulated cAMP accumulation to approximately 45% of control (Fig. 10A). However, the potency of the response was not significantly altered (EC₅₀ ± S.E.M.: control, 81 ± 58; RanBP10, 93 ± 48; RanBP9, 137 ± 12 nM). To establish whether the inhibitory effect of RanBP10 or RanBP9 was due to changes in D₁ receptor binding activity, membranes prepared from cells expressing the D₁ receptor ± RanBP10 or RanBP9 were assayed using the D₁-selective radioligand [³H]SCH23390. It is noteworthy that coexpression of RanBP10 did not significantly alter D₁ receptor expression, whereas coexpression of RanBP9 decreased D₁ receptor binding by ~50% (Fig. 10B). These results suggest that the effect of RanBP9 on D₁ receptor signaling may, at least in part, be explained by decreased receptor expression. In contrast, al-

Fig. 6. RanBP10 and RanBP9 coimmunoprecipitate with the D₁ receptor. The FLAG-D₁ receptor was expressed in HEK293T cells either alone or with RanBP10-GFP or RanBP9-GFP. Membranes were prepared and solubilized followed by immunoprecipitation of the D₁ receptor using anti-FLAG antisera as described under Materials and Methods. After resolution of the precipitates on SDS-PAGE, the gels were blotted with antisera directed to GFP (top) to detect RanB10 or RanBP9 or antisera directed to the D₁ receptor (bottom). The experiment shown is representative of three independent experiments.

Fig. 7. D₁ receptor and RanBP9 colocalization in rat kidney. D₁R (red) and RanBP9 (green) are expressed in the brush border (BB), apical membrane, and to a lesser extent observed in the cytoplasm of the proximal tubule (PT) (original magnification, 400×). The immunofluorescence images were acquired using Zeiss 510 confocal laser scanning microscope. Colocalization of the D₁ receptor and RanBP9 is indicated by the yellow color in the merged image. The experiment shown is representative of three independent experiments.
terations in receptor expression cannot explain the functional effect of RanBP10 expression on D₁ receptor signaling.

**RanBP10 and RanBP9 Increase Basal D₁ Receptor Phosphorylation by a PKC-Dependent Mechanism.** We reported previously that the D₁ receptor seems to be constitutively phosphorylated by PKC in the basal state and that this attenuates agonist-induced cAMP accumulation (Rex et al., 2008). We further showed that EtOH selectively attenuated the kinase activities of PKCγ and PKCδ, decreased basal D₁ receptor phosphorylation, and potentiated cAMP accumulation (Rex et al., 2008). Given that RanBP9 and RanBP10 associate with both PKCγ and PKCδ and with the D₁ receptor, we wondered whether RanBP9 or RanBP10 might scaffold PKCδ and PKCγ in close proximity to the D₁ receptor to facilitate receptor phosphorylation. To test this hypothesis, we examined the effect of RanBP10 and RanBP9 on D₁ receptor phosphorylation under basal or agonist-stimulated conditions using an in situ phosphorylation assay. It is noteworthy that coexpression of either RanBP10 or RanBP9 with the D₁ receptor significantly elevated basal D₁ receptor phosphorylation (Fig. 11). As expected, DA stimulation increased D₁ receptor phosphorylation compared with basal. We showed previously that the agonist-induced phosphorylation of the D₁ receptor is mediated through GRK pathways (Gardner et al., 2001; Rankin et al., 2006). It is noteworthy that in the presence of RanBP9 or RanBP10, the percentage increase in phosphorylation in response to agonist was similar to or slightly less than that observed in control cells (Fig. 12), suggesting that the main effect of RanBP9 or RanBP10 was to increase the basal phosphorylation state of the D₁ receptor.

We were particularly interested in the mechanisms responsible for the RanBP10- and RanBP9-mediated potentiation of basal D₁ receptor phosphorylation and whether or not this might be mediated by PKCs as hypothesized. To test this, we used the in situ phosphorylation assay described above except that the cells were treated with the PKC inhibitors, Gö6983 and Gö6976, before incubation in basal media. Gö6983 and Gö6976 are known to inhibit several PKC isozymes including PKCδ and PKCγ (Martiny-Baron et al., 1993; Stempka et al., 1999). As described previously (Gardner et al., 2001; Rex et al., 2008), treatment with PKC inhibitors markedly decreased basal D₁ receptor phosphorylation (Fig. 12). However, treatment with the PKC inhibitors significantly attenuated the RanBP10- and RanBP9-promoted increase in basal D₁ receptor phosphorylation, Gö6983 exhibiting a somewhat greater effect than Gö6976 (Fig. 12). Taken together, these results suggest a role for PKCs, potentially PKCδ or PKCγ, in the RanBP10- and RanBP9-dependent modulation of D₁ receptor signaling.

**Discussion**

The major finding of the current investigation is the identification of RanBP9 and RanBP10 as interacting proteins for protein kinase C isozymes γ and δ and for the D₁ dopamine receptor. Our proteomic screen initially identified RanBP10 as an interacting protein for PKCγ, although coimmunoprecipitation analyses subsequently revealed that both PKCγ and PKCδ were also capable of interacting with the structurally related protein RanBP9. Although our initial interest was in attempting to identify interacting proteins that might impart EtOH sensitivity to PKCγ/δ, neither RanBP9 nor RanBP10 seemed to affect the kinase activities of PKCγ or PKCδ. Rather, overexpression of either RanBP protein actually blocked the EtOH-mediated inhibition of PKCγ/δ activities. Although other explanations are possible, it is tempting to speculate that this effect could be due to RanBP9 or RanBP10 preventing the interaction of the PKCs with another interacting protein that might be involved in the EtOH modulation.

**Fig. 8.** Coimmunoprecipitation of the D₁ receptor and RanBP9 endogenously expressed in kidney tissue. Immortalized renal proximal tubule cells (passage, ~20, 80% confluent), made quiescent by incubating the cells for 2 h in medium without FBS, were treated with vehicle (Veh; sterile water, 10 min, lane 1) or a D₁-like agonist, fenoldopam (Fen; 1 μM, 10 min, lane 2). Lysates of treated cells or IgG were immunoprecipitated (IP) and immunoblotted (IB) as shown. To determine the specificity of the bands, normal rabbit IgG was used for immunoprecipitation and served as the negative control (lane 3), and Veh-cell lysate (immunoblot, lane 4) was used as positive control. The studies were performed three times with similar results.

**Fig. 9.** Cellular colocalization of the D₁ receptor and RanBP9 in brain. Brain sections through the striatum from transgenic mice in which D₁ receptors are marked with EGFP were processed for fluorescence immunohistochemical localization of RanBP9 (top) and RanBP10 (bottom). Immunoreactivity for both RanBP9 (RanBP-IR) and RanBP10 (RanBP-IR) colocalize with both D1-EGFP-positive striatal neurons (yellow arrows) and D1-EGFP-negative striatal neurons (blue arrows). Thus, both RanBP9 and RanBP10 seem to colocalize with striatal neurons expressing the D₁ receptor.
Because our overall goal was the investigation of PKC-mediated regulation of D1 receptor signaling, we wondered whether either PKC-interacting protein, RanBP9 or RanBP10, could also interact with the D1 receptor and affect its function. Our coimmunoprecipitation analyses revealed that both RanBP proteins were indeed capable of interacting with D1 receptor complexes when expressed in HEK293 cells. Moreover, RanBP9 and the D1 receptor are endogenously expressed in the proximal tubule of the rat kidney, and RanBP9 and the D1 receptor were found to coimmunoprecipitate using primary cultures of human renal proximal tubule cells. We also found that both RanBP9 and RanBP10 and the D1 receptor were partially colocalized at the cellular level within brain tissue.

Functionally, the association of RanBP9 and RanBP10 with the D1 receptor seems to attenuate signaling. This was evident as a decrease in maximal cAMP accumulation elicited by receptor stimulation. It is noteworthy that overexpression of RanBP9 attenuated D1 receptor expression, whereas RanBP10 did not. This might reflect fundamental differences in the interactions of RanBP9 and RanBP10 with the receptor that will merit further investigation. Although a decrease in receptor expression by RanBP9 could contribute to the attenuated cAMP accumulation, this cannot explain the functional inhibition by RanBP10, suggesting that other mechanisms must be involved. It is noteworthy that overexpression of both RanBP9 and RanBP10 led to an increase in the basal phosphorylation state of the D1 receptor. We found previously that the D1 receptor is constitutively phosphorylated by PKC and that this attenuates receptor signaling (Rex et al., 2008). Based on our findings that RanBP9 and RanBP10 associate with PKCδ and PKCγ, we examined the possibility that the increase in basal D1 receptor phosphorylation was mediated by PKC. Indeed, HEK293T cells pretreated with select PKC inhibitors blocked the RanBP-dependent increase in basal D1 receptor phosphorylation. Taken together, we propose that RanBP9 and RanBP10 regulate D1 receptor phosphorylation and signaling through a PKC-dependent mechanism.

The identification of RanBP9 and RanBP10 as dual interacting proteins for the D1 receptor and PKCs is consistent with their function as scaffolding proteins. Both RanBP9 and RanBP10 contain multiple functional domains, and there is a growing body of evidence that they both function as scaffolding molecules in immune and neural tissues (Murrin and Talbot, 2007). Of particular interest are RanBP9's interactions with membrane receptors, especially those within the GPCR superfamily such as the μ-opioid receptor (MOR) and mGluRs. RanBP9 associates with the Gαi-coupled MOR and mGlur1 (Murrin and Talbot, 2007). Overexpression of RanBP9 reduces MOR-stimulated ERK activation and inhibits β-arrestin-mediated receptor internalization (Talbot et al., 2009). However, in contrast to our findings with the D1 receptor, RanBP9 does not alter MOR phosphorylation or affect its regulation of cAMP production (Talbot et al., 2009). RanBP9 also associates with mGluR2 and mGluR8 in both HEK293 cells and the synaptic layers of the retina (Seebahn et al., 2008), although the role of RanBP9 in regulating mGluR2 and mGluR8 remains to be elucidated.

![Fig. 10. Effect of RanBP10 and RanBP9 on D1 receptor expression and signaling. A, RanBP10 or RanBP9 was co-expressed with the D1 receptor in HEK293T cells followed by assessment of DA-stimulated cAMP accumulation. The curves on the left are representative experiments, whereas the histograms on the right represent averaged data from three experiments using 100 μM DA as the stimulus. Expression of either RanBP10 or RanBP9 decreased maximal DA-stimulated cAMP accumulation compared with control (D1 receptor + empty vector): Emax ± S.E.M. values for D1R + RanBP10 and D1R + RanBP9 were 47 ± 6 and 46 ± 10% of control, respectively. B, saturation radioligand binding experiments were performed on membranes prepared from control cells (D1 receptor + empty vector) or cells expressing the D1 receptor along with either RanBP10 or RanBP9. The curves on the left are representative experiments, whereas the histograms on the right represent averaged Bmax data from three experiments. RanBP9 coexpression reduced the D1 receptor Bmax value to 60 ± 14% of control. (*, p < 0.05, **, p < 0.005, paired Student’s t test).](molpharm.aspetjournals.org@ASPET Journals on June 23, 2017)
lating glutamatergic signaling is currently unclear (Seebahn et al., 2008). Compared with RanBP9, there is little published information regarding RanBP10-dependent modulation of signaling; however, both RanBP10 and RanBP9 are described as scaffolding proteins for MET, a membrane-bound receptor tyrosine kinase for hepatocyte growth factor (Wang et al., 2004). Association of RanBP9 with MET increases recruitment of the protein SOS to MET and enhances ERK signaling. RanBP10 also associates with MET but functions as a dominant-negative to reduce SOS recruitment and decrease ERK-mediated signaling (Wang et al., 2004).

At the cellular level, we show that RanBP9 and RanBP10 can associate with the membrane-bound D1 receptor and PKCδ/γ. These findings are consistent with the site of action of the D1 receptor and regulation by PKCδ and PKCγ. It is noteworthy that Denti et al., (2004) showed that endogenous levels of RanBP9 expressed in the epithelial cells of lung, kidney, and breast are localized to the plasma membrane and cytoplasm. In the retina, RanBP10 is localized to the synaptic processes of cholinergic amacrine cells (Seebahn et al., 2008). In contrast, RanBP10 is expressed in multiple cellular locations, including the cytoplasm and nucleus of megakaryocyte cells (Schulze et al., 2008); however, there is currently no information regarding the subcellular localization of RanBP10 in other cell types, including neurons. Collectively, the subcellular locations of RanBP9 and RanBP10 seem to depend largely on the cell type.

Consistent with the notion of RanBP9 and RanBP10 functioning as scaffolding proteins, neither RanBP protein seems to modulate the kinase activities of PKCδ or PKCγ, but rather may play a role in the spatial and temporal organization of the D1 receptor-PKC signaling complex. Several PKC-interacting proteins have been described that anchor specific PKC isozymes to the appropriate signaling complex at the correct intracellular location. For example, receptor for activated C kinase 1 (RACK1) preferentially binds to PKCβ and PKCε (Ron et al., 1994; Stebbins and Mochly-Rosen, 2001; Besson et al., 2002). RACK1 is believed to stabilize the active conformation of PKCε, and shuttle the enzyme to the correct subcellular site (Ron et al., 1999). Likewise, RACK2 associates with PKCs and translocates the kinase to the Golgi membranes (Csukai et al., 1997). Protein interacting with C kinase 1 (PICK1) is a scaffolding protein that associates with presynaptic mGluR7 and PKC. It is noteworthy that PICK1 is required for the PKC-dependent phosphorylation of mGluR7 and for stable receptor expression at the cell surface (Suh et al., 2008). It is noteworthy that mGluR7-dependent neuronal plasticity is impaired in mice lacking PICK1 (Suh et al., 2008).

In summary, we report RanBP10 and RanBP9 as novel scaffolding proteins for the D1 receptors PKCδ and PKCγ. We speculate that these dual-specificity scaffolding proteins may function as signaling integrators and dictate the efficient regulation of D1 receptor signaling by PKCδ and PKCγ.

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![Fig. 11. RanBP10 and RanBP9 increase basal D1 receptor phosphorylation. In situ phosphorylation experiments were performed on HEK293T cells expressing the D1 receptor, D1 receptor + RanBP10, and D1 receptor + RanBP9 as described under Materials and Methods. Cells were incubated with media (control) or DA (10 μM) for 10 min. Top, autoradiogram of D1 receptor immunoprecipitates from a representative in situ phosphorylation assay. The lanes in the gel correspond to the bars at the bottom. Bottom, average values of band densities for each condition. The data are normalized as the percentage of control for each individual experiment. The histograms represent the mean ± S.E.M. from four independent experiments (*, p < 0.05, **, p < 0.01, ***, p < 0.001, analysis of variance followed by Bonferroni pair-wise comparisons).

![Fig. 12. The RanBP9 and RanBP10-induced increase of D1 receptor phosphorylation is blocked by PKC inhibitors. HEK293T cells expressing the D1 receptor, D1 receptor + RanBP10, and D1 receptor + RanBP9 were treated with the PKC inhibitors Go6983 (10 μM) or Go6976 (10 μM) for 60 min followed by the assessment of basal phosphorylation using an in situ phosphorylation assay as described in Fig. 11. Top, autoradiogram of D1 receptor immunoprecipitates from a representative in situ phosphorylation assay. The lanes in the gel correspond to the bars at the bottom. Bottom, average values of band densities for each condition. The data are normalized as the percentage of control for each individual experiment. The histograms represent the mean ± S.E.M. from three independent experiments (analysis of variance followed by Bonferroni pair-wise comparisons; *, p < 0.05, **, p = 0.056).
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References


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