Specificity of Collybistin-Phosphoinositide Interactions

**IMPACT OF THE INDIVIDUAL PROTEIN DOMAINS**

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Michaela Ludolphs†, Daniela Schneeberger‡, Tolga Soykan‡, Jonas Schäfer†, Theofilos Papadopoulos**, Nils Brose†, Hermann Schindelin†, and Claudia Steinem††

From the †Institute of Organic and Biomolecular Chemistry, University of Göttingen, Tamannstrasse 2, 37077 Göttingen, Germany, ‡Rudolf Virchow Center for Experimental Biomedicine, University of Würzburg, Josef-Schneider-Strasse 2, 97080 Würzburg, Germany, §Department of Molecular Neurobiology, Max Planck Institute for Experimental Medicine, Hermann-Rein-Strasse 3, 37075 Göttingen, Germany, and **Universitätsmedizin Göttingen, Department of Molecular Biology, Humboldtallee 23, 37073 Göttingen, Germany

The regulatory protein collybistin (CB) recruits the receptor-scaffolding protein gephyrin to mammalian inhibitory glycinergic and GABAergic postsynaptic membranes in nerve cells. CB is tethered to the membrane via phosphoinositides. We developed an in vitro assay based on solid-supported 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine membranes doped with different phosphoinositides on silicon/silicon dioxide substrates to quantify the binding of various CB2 constructs using reflectometric interference spectroscopy. Based on adsorption isotherms, we obtained dissociation constants and binding capacities of the membranes. Our results show that full-length CB2 harboring the N-terminal Src homology 3 (SH3) domain (CB2_{SH3+}) adopts a closed and autoinhibited conformation that largely prevents membrane binding. This autoinhibition is relieved upon introduction of the W24A/E262A mutation, which conformationally “opens” CB2_{SH3+} and allows the pleckstrin homology domain to properly bind lipids depending on the phosphoinositide species with a preference for phosphatidylinositol 3-monophosphate and phosphatidylinositol 4-monophosphate. This type of membrane tethering under the control of the release of the SH3 domain of CB is essential for regulating gephyrin clustering.

The function of neuronal synapses and the dynamic regulation of their efficacy depend on the assembly of the postsynaptic neurotransmitter receptor apparatus. The main scaffolding protein of inhibitory glycinergic and GABAergic postsynapses in mammals is gephyrin (1, 2), whose recruitment to the postsynaptic membrane is controlled by the adaptor protein collybistin (CB) (3). Loss of CB results in a strong reduction of gephyrin and GABA_A receptor clusters in several regions of the forebrain, which demonstrates the essential role of CB in the assembly and maintenance of GABAergic postsynaptic structures (4).

CB belongs to the Dbl family of guanine nucleotide exchange factors. In mouse, four differently spliced CB mRNAs are present (CB1_{SH3+}, CB2_{SH3+}, CB3_{SH3+}, and CB3_{SH3+}). All four mRNAs encode a Dbl homology (DH) and a pleckstrin homology (PH) domain. The three major variants (CB1_{SH3+}, CB2_{SH3+}, and CB3_{SH3+}) encode CBs with an additional N-terminal Src homology 3 (SH3) domain but differ in their C termini. A fourth variant (CB2_{SH3−}) encodes a CB2 isoform lacking the SH3 domain (Fig. 1) but is very rare (5) as its protein product is not detectable in mouse brain (6). Importantly, the PH domain of the different CBs is required for proper function as indicated by the fact that its deletion abolishes the plasma membrane targeting of gephyrin-CB complexes when cotransfected in HEK293 cells and causes a loss of dendritic gephyrin clusters in dissociated rat cortical neurons (5).

In vitro binding studies utilizing a variety of inositol headgroups, soluble phosphoinositide analogs, and liposomes containing phosphoinositides showed that PH domains bind phosphoinositides with a broad range of selectivity and affinity (7–10). An early membrane activation model suggested that the PH domain of CB binds to phosphatidylinositol 3,4,5-trisphosphate (PI(3,4,5)P_3) (1). In contrast, experiments with immobilized phosphoinositides and purified glutathione S-transferase (GST)-tagged CB variants in overlay assays indicated that the PH domain of CB specifically binds phosphatidylinositol 3-monophosphate (PI(3)P) (11, 12), and subsequent studies verified the binding of CB to PI(3)P (6, 13). However, most of the relevant experiments were conducted with lipids spotted on blotting membranes, which have been shown to be less reliable than other techniques (14). Hence, the question arises as to whether the phosphoinositide specificity of CB observed with overlay assays properly reflects the lipid binding specificity of CB in intact phospholipid bilayers. That is a critical issue is...
were grown to an approximately required for GABAergic synapse formation only under specific stimulatory conditions (16). Instead, PI(3,4)P2, PI(4,5)P2, and PI(3,4,5)P3 are primarily localized to the plasma membrane; PI(4)P is enriched in the Golgi complex and present at the plasma membrane; and PI(3,5)P2 is found in compartments of the late endosomal pathway (Fig. 2) (17, 18).

In the present study, we assessed the lipid binding specificity of CB in a more natural phospholipid bilayer environment that reflects the situation in live cells more closely. For that purpose, we used fluid planar lipid bilayers composed of 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) doped with different phosphoinositides. Immobilized on a silicon/silicon dioxide substrate, these membranes enabled us to monitor the specific interaction of CB variants with receptor lipids in a time-resolved and label-free manner by means of reflectometric interference spectroscopy (RIfS) (Fig. 3A) (19–21). RIfS is a well established technique to quantitatively monitor protein-receptor interactions at solid-supported membranes (22–25).

**Experimental Procedures**

**Materials**—Phosphoinositides (PI(3)P, PI(4)P, PI(3,4)P2, PI(3,5)P2, PI(4,5)P2, and PI(3,4,5)P3) were obtained as C16 derivatives from Echelon Biosciences (Salt Lake City, UT). POPC, 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-L-serine (POPS), L-α-phosphatidylinositol (PI) from soy and 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-(1’-racemic glycero) (POPG) were purchased from Avanti Polar Lipids (Alabaster, AL). BL21(DE3) competent *Escherichia coli* cells were from Invitrogen. Chitin resin was obtained from New England Biolabs (Ipswich, MA). Silicon wafers were purchased from Silicon Materials (Kaufierung, Germany).

**Protein Purification**—The CB variants CB2PH, CB2SH3, and CB2SH3/W24A/E262A were obtained by recombinant expression as described previously (6). Briefly, plasmids based on the IMPACT system vector pTXB1 (CB2SH3 and CB2SH3/W24A/E262A) or pTYB12 (CB2PH and CB2SH3) encoding CB2 isoforms with an N-/C-terminal intein tag were transformed into *E. coli* BL21(DE3) cells. Cells were grown to an *A*\textsubscript{600} of 0.8–1.0 at 37 °C in LB-Miller medium. Protein expression was induced by addition of 0.5 mM isopropyl-β-D-thiogalactopyranoside. After overnight growth at 15 °C, cells were harvested by centrifugation (4,500 × g, 20 min, 4 °C). Cells were resuspended in lysis buffer (250 mM NaCl, 20 mM HEPES, 2 mM EDTA, 10% (v/v) glycerol, pH 8.0), and cell lysis was performed using a microfluidizer (1 kbar, three cycles, iced-cooled; LM 10 processor, Microfluidics, Westwood, MA). After centrifugation (70,000 × g, 30 min, 4 °C), the supernatant was applied to the equilibrated chitin resin for 1 h. The resin was rinsed with >1 liter of washing buffer (1 M NaCl, 20 mM HEPES, 2 mM EDTA, pH 8.0), and protein cleavage was induced by incubation with 50 mM dithiothreitol (DTT) in buffer G (250 mM NaCl, 20 mM HEPES, 2 mM EDTA, pH 8.0) for >40 h. After elution with buffer G containing 5 mM DTT, the protein solution was diluted to 50 mM NaCl. The protein was purified using a Mono Q 10/100GL column (GE Healthcare) with a linear NaCl gradient from 50 mM to 1 M NaCl in buffer (20 mM HEPES, 2 mM EDTA, 2 mM β-mercaptoethanol, pH 8.0). Proteins were stored at 4 °C. In the case of CB2SH3, 20 mM Tris instead of HEPES was used in all buffers. Concentration and dialysis to buffer A (100 mM NaCl, 25 mM HEPES, 0.5 mM DTT, 0.5 mM EDTA, pH 7.2) was performed by ultrafiltration using spin concentrators (Sartorius, Göttingen, Germany). Protein concentrations were determined by UV/visible spectroscopy using extinction coefficients of $\epsilon_{280} = 37,930 \text{ M}^{-1} \text{ cm}^{-1}$ (CB2PH), $\epsilon_{280} = 70,000 \text{ M}^{-1} \text{ cm}^{-1}$ (CB2SH3), $\epsilon_{280} = 98,945 \text{ M}^{-1} \text{ cm}^{-1}$ (CB2SH3), or $\epsilon_{280} = 93,445 \text{ M}^{-1} \text{ cm}^{-1}$ (CB2SH3/W24A/E262A).

**Vesicle Preparation**—Stock solutions of POPC, POPS, POPG, and PI were prepared in chloroform, whereas the phosphoinositides were dissolved in a mixture of chloroform/methanol/water according to the manufacturer’s instructions. Lipid stock solutions were added to chloroform to obtain the desired lipid molar ratios, and the organic solvents were evaporated with a gentle nitrogen stream at 30 °C. Lipid films were further dried under vacuum for at least 3 h at 32 °C. For the preparation of small unilamellar vesicles (SUVs), the lipid film was rehy-
drated by incubation with 500 μl of buffer for 20 min followed by vortexing for 30 s (three cycles, every 5 min). The vesicle suspension was then sonicated using an ultrasonic homogenizer (30 min, 65%, four cycles; Sonopuls HD2070, Bandelin, Berlin, Germany) to generate SUVs.

Reflectometric Interference Spectroscopy—RIfS was used to monitor the formation of supported lipid bilayers and subsequent protein binding in a time-resolved manner (26). The experimental setup was described in detail previously (Fig. 3A) (22). Reflectometric interference spectra were monitored using a NanoCalc-2000 visible/near-IR spectrometer (Ocean Optics, Dunedin, FL). Spectra were recorded every 2 s and were analyzed using a MATLAB routine. To provide a substrate for interference fringes, silicon wafers with a 5-μm-thick SiO2 layer were used. Silicon substrates (1 × 2 cm2) were rinsed with ethanol and water. To hydrophilize the silicon substrates, they were incubated in an aqueous solution of NH3 and H2O2 (H2O/NH3/H2O2, 5:1:1) for 20 min at 70 °C followed by treatment with oxygen plasma (30 s, 50% power). Afterward, the hydrophilic substrate was rinsed with buffer, and SUVs were added. After the spreading process of the SUVs was finished, as indicated by a constant optical thickness, the remaining lipid material was removed by rinsing with buffer A. To prevent nonspecific protein binding, the measuring chamber was flushed with a bovine serum albumin-containing buffer solution (1 mg/ml in buffer A) and rinsed again with buffer A.

CD Spectroscopy—CD spectra were used to determine the secondary structure elements of the isolated proteins and to compare them with the crystal structure of CB2SH3 (Protein Data Bank code 4MT6) (6). CD spectra (λ = 190 – 260 nm) were recorded at 20 °C with a J-1500 spectrometer (Jasco, Gross-Umstadt, Germany) equipped with a cuvette (d = 0.1 mm). Five spectra were averaged, background was subtracted (buffer solution), and the mean residue ellipticity was calculated. Secondary structure analysis was performed with the program DichroWeb using the CDSSTR method and reference data set 4 (27). Table 1 summarizes the fractions of the secondary structure elements as obtained from data evaluation using DichroWeb.

Results

Formation of Supported Lipid Bilayers—To obtain planar supported lipid bilayers, SUVs were spread onto the silicon/silicon dioxide surface (Fig. 3B). To obtain membranes with high surface coverage, optimum spreading conditions of the vesicles to form planar bilayers are required. In a previous study, we showed that SUVs composed of POPC doped with PI(4,5)P2 do not spread to a continuous bilayer on silicon/silicon dioxide substrates at pH 7.4 if the PI(4,5)P2 concentration is larger than 4 mol % (23). However, if the pH is reduced to 4.8, the net negative charge of PI(4,5)P2 and the negative surface charge of the silicon substrate are reduced (28, 29). In the present study, we doped POPC membranes with 10 mol % of the different phosphatidylinositol phosphates (PIPs), namely PI(3)P, PI(4)P, PI(3,4)P2, PI(3,5)P2, PI(4,5)P2, and PI(3,4,5)P3, 10 mol % of the potential receptor lipid was chosen to ensure a good signal-to-noise ratio. As the PIPs vary in their phosphorylation pattern and thus in their overall net negative charge, the pH value of the buffer was adjusted accordingly. For SUVs containing 10 mol % PI(3)P and PI(4)P, a buffer at pH 6.4 was used (20 mM citrate, 50 mM KCl, 0.1 mM EDTA, 0.1 mM NaN3, pH 6.4), whereas for SUVs composed of POPC and 10 mol % PI(3,4)P2, PI(3,5)P2, PI(4,5)P2, or PI(3,4,5)P3, a citrate buffer at pH 4.8 (20 mM citrate, 50 mM KCl, 0.1 mM EDTA, 0.1 mM NaN3, pH 4.8) was chosen. SUVs containing POPC only, POPC/POPS (8:2), POPC/POPG (9:1 and 8:2), or POPC/PI (9:1) were prepared and spread in buffer A, which was also used for protein binding studies.

The spreading process of the SUVs was monitored in a time-resolved manner by means of RIfS. A characteristic time trace is depicted in Fig. 4A. From the maximum change in optical thickness (ΔOT), the quality of the resulting bilayer can be assessed. After the spreading process was completed, the system was rinsed with buffer A. No changes in bilayer thickness were observed, which is indicative of a stable planar solid-supported bilayer (Fig. 4A).

From the ΔOT values at saturation, the physical thickness of the bilayer (d membrane) was calculated using ΔOT =
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**FIGURE 4. Membrane preparations analyzed by RIfS.** A, time-resolved change in membrane thickness $d_{\text{membrane}}$ during the spreading of SUVs composed of POPC/PI(3,4,5)P$_3$ (9:1) on a silicon wafer. a, addition of SUVs (0.4 mg/ml) in 20 mM citrate buffer, pH 4.8; b, rinsing with buffer. B, membrane thicknesses $d_{\text{membrane}}$ for different lipid compositions. Error bars show the S.D. of the mean obtained from at least nine independent membrane preparations.

$n_{\text{lipid}}d_{\text{membrane}}$ with a refractive index of $n_{\text{lipid}} = 1.5$. For POPC membranes, a bilayer thickness of $d_{\text{membrane}} = 4.28 \pm 0.04$ nm was obtained (Fig. 4B). This value is in good agreement with measurements using small angle neutron and x-ray scattering ($d_{\text{membrane}} = 3.98 \pm 0.08$ nm) (30). For POPC membranes containing POPG (9:1 and 8:2), membrane thicknesses of 4.18 ± 0.04 and 4.02 ± 0.01 nm were obtained, whereas for POPC/POPS (8:2) membranes, a thickness of 4.1 ± 0.2 nm was measured; these values are in good agreement with the thickness of a pure POPC bilayer and thus indicate rather defect-free bilayers. For membranes containing PI, PI(3)P, and PI(4)P, slightly thinner bilayers were obtained (3.8 ± 0.4 nm for PI(3)P, 3.7 ± 0.7 nm for PI(4)P, and 3.5 ± 0.4 nm for PI-containing bilayers; Fig. 4B), which indicates that some defects are present in these bilayers. Compared with pure POPC bilayers, a membrane surface coverage of at least 90% was still obtained. To prevent nonspecific protein binding to these defects, the membranes were incubated with a bovine serum albumin (BSA) solution prior to the addition of the CB variant as described under “Experimental Procedures.” The slightly lower membrane surface coverage is probably a result of the repulsion between the negatively charged vesicles and the negatively charged silicon dioxide surface at the given pH (23) but is still fully sufficient for the experiments given the fact that the free surface area is blocked with BSA. Membranes doped with PI(3,4)P$_2$, PI(3,5)P$_2$, PI(4,5)P$_2$, or PI(3,4,5)P$_3$ show slightly larger membrane thicknesses as compared with POPC-only membranes (Fig. 4B), which might be explained by the strongly phosphorylated inositol headgroups that protrude about 0.6 nm from the membrane surface (31).

**Secondary Structure Analysis of CB Variants—**Circular dichroism spectra were taken and analyzed for all CB variants. The fraction of the secondary structure elements was compared with the crystal structure to provide evidence for proper protein folding under the chosen buffer conditions (Table 1). The crystal structure of CB2$_{\text{PH}}$ shows nine antiparallel $\beta$-strands and a C-terminal $\alpha$-helix, which is reflected in 39% $\beta$-strands and 14% $\alpha$-helices. CB2$_{\text{SH3-}}$ consists of a DH/PH tandem domain with a DH domain formed only by $\alpha$-helices. This is reproduced in the large increase (54%) in $\alpha$-helical content of CB2$_{\text{SH3-}}$. Full-length CB2$_{\text{SH3+}}$ is composed of the DH/PH tandem domain and the N-terminal SH3 domain. The SH3 domain consists of five short $\beta$-strands connected by unordered loop regions, leading to an increase in the $\beta$-strand fraction (23%) as compared with CB2$_{\text{SH3-}}$. In contrast to the crystal structure, we found a larger $\alpha$-helical and larger $\beta$-strand fraction at the expense of random coil. This can be explained by the fact that the linker between the SH3 and DH domain is missing in the crystal structure and hence does not contribute to our calculations.

**Interaction of CB2$_{\text{PH}}$ with Different Phosphoinositides—**The planar continuous lipid bilayers doped with different PIPs and prepared as described above are a prerequisite for the analysis of the specificity of CB binding. As binding to phosphoinositides is known to be mediated by a region of positively charged amino acids in the PH domain of CB (32), we started out by studying the isolated PH domain of CB (CB$_{\text{PH}}$) and its specificity for certain PIPs. Protein concentration-dependent binding studies were performed by means of the RIfS setup. Fig. 5A shows a typical result of a RIfS experiment. Adding CB2$_{\text{PH}}$ to a membrane composed of POPC doped with 10 mol % of PI(3)P results in a change in thickness due to interactions between the protein and the membrane-embedded lipids. With increasing protein concentration, the thickness of the protein layer increases until the maximum protein layer thickness ($d_{\text{protein-max}}$) is reached. Rinsing with buffer A leads to a partial decrease in thickness, indicating that about 50% of the protein is reversibly bound and desorbs upon rinsing. It is a general phenomenon that proteins bound via a receptor lipid to a planar membrane only partially desorb from the membrane upon rinsing (23, 33). This can be explained as follows. (i) Proteins that bind in a one-to-one fashion to the receptor lipid desorb from the membrane upon rinsing with the corresponding rate constant of desorption, $k_{\text{off}}$. However, part of the proteins might bind more than one receptor lipid or bind additionally to other lipid headgroups so that they do not desorb upon rinsing. (ii) If the membrane-bound proteins laterally interact with each other and form protein clusters, this second interaction on the membrane surface prevents the proteins from desorption upon rinsing.

From the saturation values reached after each addition of the respective protein, $d_{\text{protein-max}}$ was extracted (see Fig. 5A, inset) and...
**TABLE 1**

Secondary structure elements of the different CB variants used in this study in comparison with the fractions calculated (Calc.) from the crystal structure of CB2\textsubscript{SH3+} (Protein Data Bank code 4MT6) (6)

| Structure          | CB2\textsubscript{PH} | Calc. | CB2\textsubscript{SH3} | Calc. | CB2\textsubscript{SH3+} | Calc. | CB2\textsubscript{SH3+}/W24A/E262A | Calc. |
|--------------------|------------------------|-------|-------------------------|-------|-------------------------|-------|-------------------------------|-------|
| α-Helix            | 0.14                   | 0.18  | 0.54                    | 0.44  | 0.51                    | 0.53  | 0.39                          |       |
| β-Strand           | 0.39                   | 0.31  | 0.09                    | 0.14  | 0.23                    | 0.11  | 0.15                          |       |
| β-Turn             | 0.17                   | 0.06  | 0.13                    | 0.07  | 0.07                    | 0.14  | 0.06                          |       |
| Random coil        | 0.29                   | 0.45  | 0.23                    | 0.35  | 0.18                    | 0.22  | 0.40                          |       |

We are aware of the fact that the Langmuir model is only valid in the case of fully reversible protein binding, and more elaborate models are required to account for the irreversibility (33–35). However, as the origin of the irreversibility is as yet not known, we refrained from a more sophisticated model and used the obtained $K_D$ values to compare the affinities among the different PIPs. Control experiments with POPC-only membranes demonstrated that binding of CB2\textsubscript{PH} is specific; i.e., no interaction of the protein with POPC membranes lacking phosphoinositides was observed (data not shown). Also no protein binding was observed on membranes composed of POPC/POPS (8:2), indicating that a mere negative surface charge density is not sufficient for CB2\textsubscript{PH} to bind.

We obtained adsorption isotherms for different PIPs (Fig. 6) from which the dissociation constants and protein layer thicknesses were determined (Table 2). The isotherms indicate that CB2\textsubscript{PH} does not significantly bind to PI(3,4)P\textsubscript{2} as indicated by the maximum change in protein layer thickness of only 0.2–0.3 nm. These values are too low to be attributed to specific binding. Thus, all measurements with such low protein surface coverage were not subjected to a determination of dissociation constants.

In contrast to this low nonspecific binding, CB2\textsubscript{PH} binds to all other PIPs with a dissociation constant in the 0.1 μM range with a slight preference for monophosphorylated PIPs. The dissociation constants for the diphosphorylated PI derivatives PI(3,5)P\textsubscript{2} and PI(4,5)P\textsubscript{2} and the triphosphorylated PI(3,4,5)P\textsubscript{3} are about 4–8 times larger. The largest protein layer thickness was found for PI(3,4,5)P\textsubscript{3}-doped membranes with $d_{\text{protein-max}} = 2.8 \pm 0.3$ nm. Assuming a maximum protein surface coverage of 56% according to the scaled particle theory (25) and a diameter of the PH domain of 4.2 nm as obtained from the crystal structure (12), one would expect a protein layer thickness of 2.4 nm. Comparing the theoretical and experimental values, we conclude that full protein coverage is reached in the case of PI(3,4,5)P\textsubscript{3}, whereas for the other PIPs, a lower surface coverage is found.

Interaction of CB2\textsubscript{SH3} with Different Phosphoinositides—It has been proposed that CB exists in a closed and autoinhibited conformation in which the SH3 domain interacts with the DH and PH domains, thus preventing its membrane recruitment (6). Hence, for CB2 lacking the SH3 domain, it is expected that it preferentially binds to phosphoinositides, whereas the full-length protein interacts only weakly. In light of this hypothesis, we first studied the binding of CB2\textsubscript{SH3+}, which harbors the DH domain and the PH domain but lacks the SH3 domain, to PIP-containing membranes. Again, adsorption isotherms were monitored for different PIPs, and the dissociation constants as well as the maximum protein layer thicknesses were determined by fitting a Langmuir adsorption isotherm to the data (Fig. 6). As expected, CB2\textsubscript{SH3} binds with submicromolar affinity to all PIPs except for PI(3,4)P\textsubscript{2}, demonstrating that the PH domain of CB is accessible for PIP binding. The same trend was observed for the di- and triphosphorylated PIPs with slightly larger $K_D$ values (Fig. 6 and Table 2). Due to the size of the additional DH domain, the maximum protein layer thickness on the membrane increased as expected except for PI(3,5)P\textsubscript{2}, which also shows the smallest binding affinity among the different PIPs (Fig. 6). The largest maximum protein layer thicknesses of 3.5–4 nm were determined for PI(4,5)P\textsubscript{2} and PI(3,4,5)P\textsubscript{3}, similar to the results obtained for CB2\textsubscript{PH}. Based on the protein dimensions obtained from the crystal structure of CB2\textsubscript{PH} to PI(3)P. A, characteristic time trace of a RIFS experiment showing the specific binding of CB2\textsubscript{PH} to a PI(3)P-containing membrane. Different concentrations of CB2\textsubscript{PH} were added at time points as indicated by arrows (a, 0.038 μM; b, 0.11 μM; c, 0.19 μM; d, 0.26 μM; e, 0.37 μM; f, 0.74 μM). After rinsing the system with buffer A (g), a fraction of the bound protein desorbs from the membrane, indicating reversible binding. The inset shows the determination of the saturation value $d_{\text{protein}}$ (Eq. 1).

$$d_{\text{protein}} = d_{\text{protein-max}} \frac{c}{K_D + c} \quad (\text{Eq. 1})$$

The solid gray line shows the resulting adsorption isotherm. The dissociation constant $K_D$ as well as $d_{\text{protein-max}}$, which reflects the binding capacity of the membrane, were determined by fitting a simple Langmuir isotherm (Equation 1) to the data with $KD$ values (Fig. 6 and Table 2). Due to the size of the DH and PH domains, the largest maximum protein layer thickness was found for PI(3,4,5)P\textsubscript{3}-doped membranes with $d_{\text{protein-max}} = 2.8 \pm 0.3$ nm. Assuming a maximum protein surface coverage of 56% according to the scaled particle theory (25) and a diameter of the PH domain of 4.2 nm as obtained from the crystal structure (12), one would expect a protein layer thickness of 2.4 nm. Comparing the theoretical and experimental values, we conclude that full protein coverage is reached in the case of PI(3,4,5)P\textsubscript{3}, whereas for the other PIPs, a lower surface coverage is found.
CB2SH3−, a theoretical protein layer thickness of 4.4 nm was estimated.

We next tested the impact of the chemical nature of the receptor lipid on the binding affinity of CB2SH3−. In accordance with the results obtained for CB2PH, no binding to negatively charged POPC/POPS (8:2) membranes was found, supporting the notion that a mere electrostatic interaction of the protein with the membrane is not responsible for CB binding. However, CB2SH3 binds to lipid membranes composed of POPC/PI (9:1) with a dissociation constant (KD = 0.14 ± 0.01 µM) similar to those of the phosphorylated PI derivatives but with a smaller protein layer thickness of dprotein-max = 0.8 ± 0.2 nm. This observation is compatible with the idea that the OH groups of the inositol headgroup together with the phosphate group at the glycerol backbone might be involved in the binding specificity of the protein.

Binding of CB2SH3+—In view of the notion that CB harboring an SH3 domain forms a closed and autoinhibited conformation where the SH3 domain interacts with the DH and PH domains, we further analyzed the influence of the SH3 domain on the binding behavior of CB by studying CB2SH3+ binding to PIP-doped membranes. If CB2SH3+ bound to PIP-doped membranes, we expected to observe a larger protein layer thickness than for CB2SH3− as CB2SH3+ is larger in size than CB2SH3−. However, in contrast to this, the observed protein layer thicknesses were greatly diminished by at least a factor of 5 (Fig. 7) independently of the chosen PIP. This finding indicates that only a minor fraction of CB2SH3+ is still capable of binding to

![Figure 6. Binding isotherms of CB2PH (black circles) and CB2SH3 (blue circles) obtained for different lipid compositions.](image)

**TABLE 2**

| Dissociation constants (KD) for the binding of CB2PH, CB2SH3, and CB2SH3/mutant to PIP-containing solid-supported membranes |
|---|
| | PI(3)P | PI(4)P | PI(3,4)P₂ | PI(3,5)P₂ | PI(4,5)P₂ | PI(3,4,5)P₃ |
| CB2SH3+ | 0.10 ± 0.01 | 0.16 ± 0.03 | 0.8 ± 0.2 | 0.60 ± 0.06 | 0.4 ± 0.1 |
| CB2SH3− | 0.32 ± 0.08 | 0.26 ± 0.06 | 1.2 ± 0.5 | 0.63 ± 0.09 | 0.3 ± 0.1 |
| CB2SH3+/mutant | 0.27 ± 0.02 | 0.25 ± 0.04 | 0.9 ± 0.1 | 1.9 ± 0.2 |
the membrane. The majority remains in an inactive state not capable of membrane binding. This is in line with the proposed “closed” and thus autoinhibited conformation of CB2SH3/H11001 (6).

Partial Activation of CB2SH3/H11001—Point mutations in CB2SH3/H11001 (e.g. CB2SH3/H11001/W24A/E262A) that perturb the intramolecular interactions within CB2SH3/H11001 relieve autoinhibition, leading to an increase in CB-triggered gephyrin clustering in COS7 cells, CB/H11545 neurons, and neuroligin-2/H11545 neurons, indicating that the corresponding mutant adopts an open conformation with an increased capacity to bind to phosphoinositide-containing membranes (6). We tested whether CB2SH3/H11001/W24A/E262A is also capable of binding to PIP-containing membranes in vitro (Fig. 7). Indeed, a significant increase in protein layer thickness for the mutant CB2SH3/H11001/W24A/E262A compared with wild-type CB2SH3/H11001 was observed for the monophosphorylated PIPs and for PI(3,5)P₂ and PI(3,4,5)P₃. These results indicate that a larger fraction of CB2SH3/H11001/W24A/E262A is in an open conformation and capable of binding to phosphoinositides with submicromolar to micromolar affinities as compared with wild-type CB2SH3/H11001. Consistent with our previous results, no binding to PI(3,4)P₂ was found. However, for PI(4,5)P₂ (Fig. 7E), also no binding of CB2SH3/H11001/W24A/E262A was observed. Importantly, the CB2SH3/H11001/W24A/E262A variant is partially selective for certain phosphoinositides with a preference for monophosphorylated phosphoinositols (PI(3)P and PI(4)P) and for PI(3,5)P₂.

Discussion

Based on an in vitro phospholipid bilayer membrane system, the present study provides important new insights into the specificity of CB2 binding to phosphoinositides and elucidates the mechanism of autoinhibition of CB2 at the molecular level. We made use of solid-supported planar bilayers doped with different phosphoinositides to provide a membrane system that partially resembles, but at the same time simplifies, the natural situation at cellular membranes. It allowed for a quantitative analysis of the binding properties of CB2 in a label-free manner by means of reflectometric interference spectroscopy. Previous studies on the specificity of CB used pure phosphoinositides spotted on synthetic membranes (protein-lipid overlay assays) (6, 12, 13). In these assays, phosphoinositides are not embedded in a lipid membrane, and thus the headgroup positions of the phosphorylated inositols are not defined and aligned, which can alter the binding specificity (14).
Here we first tested the specificity of binding of the PH domain of CB2 to different phosphoinositides, which is of key relevance as the PH domain of CB2 is functionally essential (5, 12, 36). In general, PH domains are best known for their ability to bind phosphoinositides and to be targeted to cellular membranes (10, 37). Most lipid-binding PH domains show a preference for one or several phosphoinositides, such as the PH domain of phospholipase C-\(\beta\)1, which binds specifically to PI(4,5)P\(_2\) (38). For the isolated PH domain of CB2, the dissociation constants for the phosphoinositides tested were in the 0.1–0.8 \(\mu\)M range with an up to 8 times higher affinity for monophosphorylated PI(3)P and PI(4)P. In the literature, quite a large range of \(K_D\) values can be found for PH domain binding to phosphoinositides, ranging from about 30 \(\mu\)M in the case of the interaction of the pleckstrin PH domain with PI(4,5)P\(_2\) (7) to 10 nM values for the PH domains of phospholipase C-\(\beta\)1 and Grp1-PH (39, 40). Another important aspect in this regard is the binding capacity of the membrane as a function of different phosphoinositides. We used 10 mol % of the corresponding receptor lipid, which is roughly 10 times higher than the concentrations of phosphoinositides found in native cellular membranes and was chosen to optimize the signal-to-noise ratio in our assay. If the receptor lipids were homogeneously distributed in the POPC membranes and assuming a 1:1 stoichiometry of PIP-CB2 binding as has been shown for other PH domain-containing proteins (37, 41, 42), a lower phosphoinositide concentration should have been sufficient. However, we found that at 10 mol % the maximum protein surface is reached (43), the maximum protein layer thickness obtained by RIfS experiments is a measure of the binding capacity of the membrane. Interestingly, a large protein layer thickness for CB2\(_{PH}\) was obtained for phosphoinositides with a phosphate group at the 5-position with the largest \(d_{protein-max}\) found for the most strongly phosphorylated form, PI(3,4,5)P\(_3\). The monophosphorylated phosphoinositols showed a lower binding capacity. This might be explained by a non-homogeneous, clustered distribution of the receptor lipids prior to and after protein binding (17), respectively, resulting in more than one phosphoinositide bound to CB and would require further investigations.

The submicromolar affinity of the isolated PH domain of CB2 was also seen with the DH/PH tandem domain of CB2. The slightly lower binding affinity of CB2\(_{SH3}\) for monophosphorylated phosphoinositols as compared with the isolated PH domain of CB2 is likely due to the multiple interactions between the PH and DH domains, which influence the interactions between the DH/PH domains and membrane lipids (13, 32). The maximum protein layer thickness was largest for PI(4,5)P\(_2\) and PI(3,4,5)P\(_3\), which is similar to the results obtained for CB2\(_{PH}\) and fits to the dimensions estimated from the crystal structure of CB2\(_{SH3}\) (long axis) (6). Based on the results obtained with POPC membranes containing either phosphatidylinositol or phosphatidylserine, we conclude that the molecular recognition of the OH groups of the inositol headgroup and the specific phosphorylation state are more relevant for the specific recognition of CB than an interaction driven purely by electrostatics.

In mouse brain, CB2\(_{SH3}\) exists only in trace amounts, and the majority of CB isoforms contain an additional SH3 domain (6). This SH3 domain causes CB to adopt a closed and autoinhibited conformation in which the SH3 domain interacts with the DH/PH tandem domain (6), thus preventing its membrane recruitment (Fig. 8). To analyze the phosphoinositide binding characteristics of the biologically relevant CB variants, \(i.e.\) the ones with an N-terminal SH3 domain, we studied CB2\(_{SH3}\) and a constitutively activated variant in which the autoinhibitory effect of the SH3 domain is eliminated (CB2\(_{SH3}\)/W24A/E262A). With CB2\(_{SH3}\), the amount of bound protein is greatly
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diminished as compared with CB2\textsubscript{PH} and CB2\textsubscript{SH3–}, independently of the phosphoinositide used. Taking into account the more compact structure of CB2\textsubscript{SH3–}+ which has a smaller radius of gyration ($R_g = 26.3$ Å) than CB2\textsubscript{SH3–} ($R_g = 28.3$ Å) (6), the protein size does not explain this significant decrease in protein layer thickness. Instead, the results indicate that a large fraction of CB2\textsubscript{SH3–}+ is not capable of binding to the PIP-containing membranes. This provides key support for the notion that CB2\textsubscript{SH3–}+ adopts an autoinhibited conformation that is stabilized by contacts between the SH3 domain and the DH/PH tandem domain (Fig. 8). Furthermore, our data show that a main consequence of the SH3 domain-mediated autoinhibition of CB is an inhibition of phosphoinositide binding and membrane tethering of CB (6), whereas the guanine nucleotide exchange factor activity does not seem to be affected (6, 32, 44, 45). This is different in Asef homologue, where the SH3 domain-mediated autoinhibition of Asef affects the enzymatic guanine nucleotide exchange factor activity of the DH domain (46).

To further assess the autoinhibitory influence of the SH3 domain on phosphoinositide binding of CB2, we made use of the CB2\textsubscript{SH3–}/W24A/E262A variant in which two amino acids in the SH3-DH/PH interface are exchanged. The corresponding mutation leads to an open conformation exhibiting a larger flexibility and a more elongated protein shape in solution. In lipid overlay assays, the mutant protein binds more strongly to PI(3)P as compared with wild-type CB2\textsubscript{SH3–} albeit not as well as CB2\textsubscript{SH3–} (6). A similar behavior of CB2\textsubscript{SH3–}/W24A/E262A was observed with solid-supported POPC membranes containing PI(3)P. Although the binding capacity of CB2\textsubscript{SH3–}+ was estimated to be about 3 times lower than that of CB2\textsubscript{SH3–} (Fig. 7A), it was regained to about 90% in the case of the CB2\textsubscript{SH3–}/W24A/E262A mutant. Interestingly, this regain in activity is a function of the phosphoinositide. Although the binding capacity of the mutant for the monophosphorylated phosphoinositols and for PI(3,5)P\textsubscript{2} is large, it remains partially diminished for the other PIPs, and the mutant protein does not significantly bind to phosphoinositides with two juxtaposed phosphate groups, i.e. PI(3,4)P\textsubscript{2} and PI(4,5)P\textsubscript{2}.

The present results provide the first insights into the phosphoinositide binding specificity of CB in a phospholipid bilayer context. Of most biological relevance are the data on CB2\textsubscript{SH3–}+ and its open and disinhibited CB2\textsubscript{SH3–}/W24A/E262A variant because essentially all CB isoforms expressed in murine brain carry an N-terminal SH3 domain (6). Our corresponding data nicely corroborate the notion that CB2\textsubscript{SH3–}+ is autoinhibited with respect to phosphoinositide binding and that this autoinhibition is relieved upon introduction of the W24A/E262A mutation, which “opens” the CB2\textsubscript{SH3–}+ conformation and exposes the PH domain for proper lipid binding. In neurons expressing wild-type CB, this conformational activation of phosphoinositide binding of CB is mediated by neurogin-2 (47), TC10 (44), and neurogin-4 (48), likely along with other interactors of the SH3 and PH domains of CB, to promote CB-dependent gephyrin clustering at nascent GABAergic synapses. Furthermore, our data show that CB2\textsubscript{SH3–}+ can bind to several phosphoinositide variants but show a significant preference for PI(3)P and PI(4)P in bilayer membranes. This partial selectivity, which has been observed with other assay systems as well (6, 12, 13, 36), is also seen with the isolated PH domain of CB (CB2\textsubscript{PH}) but only to a smaller degree with the N-terminally truncated CB2\textsubscript{SH3–} variant, which lacks the SH3 domain. In view of these findings and given that neurons almost exclusively express SH3 domain-containing CB isoforms, future studies on the regulation of CB function will have to focus on PI(3)P and PI(4)P signaling toward CB.

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