Lipid Binding Regulates Synaptic Targeting of PICK1, AMPA Receptor Trafficking, and Synaptic Plasticity

Wenyeng Jin,1 Woo-Ping Ge,2 Junyu Xu,1 Mian Cao,1 Lisheng Peng,1 Wingho Yung,3 Dezhi Liao,4 Shumin Duan,4 Mingjie Zhang,3 and Jun Xia1

1Department of Biochemistry, The Hong Kong University of Science and Technology, Clear Water Bay, Kowloon, Hong Kong, China, 2Institute of Neuroscience, Shanghai Institutes of Biological Sciences, Chinese Academy of Sciences, Shanghai 200031, China, 3Department of Physiology, Faculty of Medicine, The Chinese University of Hong Kong, Shatin, Hong Kong, China, and 4Department of Neuroscience, University of Minnesota, Minneapolis, Minnesota 55455

The targeting and surface expression of membrane proteins are critical to their functions. In neurons, synaptic targeting and surface expression of AMPA-type glutamate receptors were found to be critical for synaptic plasticity such as long-term potentiation and long-term depression (LTD). PICK1 (protein interacting with C kinase 1) is a cytosolic protein that interacts with many membrane proteins, including AMPA receptors via its PDZ (postsynaptic density-95/Discs large/zona occludens-1) domain. Its interactions with membrane proteins regulate their subcellular targeting and surface expression. However, the mechanism by which PICK1 regulates protein trafficking has not been fully elucidated. Here, we show that PICK1 directly binds to lipids, mainly phosphoinositides, via its BAR (Bin/amphiphysin/Rvs) domain. Lipid binding of the PICK1 BAR domain is positively regulated by its PDZ domain and negatively regulated by its C-terminal acidic domain. Mutation of critical residues of the PICK1 BAR domain eliminates its lipid-binding capability. Lipid binding of PICK1 controls the subcellular localization of the protein, because BAR domain mutant of PICK1 has diminished synaptic targeting compared with wild-type PICK1. In addition, the BAR domain mutant of PICK1 does not cluster AMPA receptors. Moreover, wild-type PICK1 enhances synaptic targeting of AMPA receptors, whereas the BAR domain mutant of PICK1 fails to do so. The BAR domain mutant of PICK1 loses its ability to regulate surface expression of the AMPA receptors and impairs expression of LTD in hippocampal neurons. Together, our findings indicate that the lipid binding of the PICK1 BAR domain is important for its synaptic targeting, AMPA receptor trafficking, and synaptic plasticity.

Key words: PICK1; AMPA receptor; synaptic plasticity; LTD; PDZ domain; BAR domain

Introduction
Regulated insertion and internalization of AMPA-type glutamate receptors at synapses are important molecular mechanisms of synaptic plasticity, including long-term potentiation (LTP) and long-term depression (LTD) (Bredt and Nicoll, 2003; Park et al., 2004). Several lines of evidence indicate that the C termini of AMPA receptors are crucial for their synaptic targeting and surface expression (Malinow and Malenka, 2002). A number of intracellular proteins that interact with the C termini of AMPA receptors have been identified (Barry and Ziff, 2002; Song and Huganir, 2002; Henley, 2003). One such protein, PICK1 (protein interacting with C kinase 1), is a cytosolic protein containing an N-terminal PDZ [postsynaptic density-95 (PSD-95)/Discs large/zona occludens-1] domain, a central BAR (Bin/amphiphysin/Rvs) domain, and an acidic domain close to the C terminus (supplemental Fig. S1A, available at www.jneurosci.org as supplemental material). The PDZ domain of PICK1 interacts with the C termini of AMPA receptor subunits (Dev et al., 1999; Xia et al., 1999). PICK1 colocalizes with the AMPA receptors at excitatory synapses and clusters the AMPA receptors when coexpressed in heterologous cells. The interaction of PICK1 with AMPA receptors was found to regulate surface expression of the AMPA receptors in a protein kinase-dependent manner (Matsuda et al., 1999; Chung et al., 2000; Perez et al., 2001; Terashima et al., 2004). Furthermore, the binding of PICK1 to AMPA receptors was found to be critical to AMPA receptor trafficking during synaptic plasticity (Matsuda et al., 2000; Xia et al., 2000; Kim et al., 2001). In a more recent study, PICK1 knock-out mice were found to be deficient in the activity-dependent insertion of glutamate receptor 2 (GluR2)-containing AMPA receptors (Gardner et al., 2005).

In addition to its PDZ domain, PICK1 was previously predicted to have a coiled-coil domain in the middle of the protein (Xia et al., 1999). Recently, a crystallographic study of the amphiphysin BAR domain suggested that this predicted coiled-coil domain is likely part of a larger BAR domain (Peter et al., 2004). The BAR domain is found in many proteins that are involved in
protein trafficking, especially endocytosis. The BAR domain of amphiphysin can directly bind and tubulate liposomes (Takei et al., 1999; Peter et al., 2004). In addition, amphiphysin was also found to interact with adaptor protein 2 (AP2), clathrin, dynamin, and syntaxin and to be critically involved in the endocytosis of synaptic vesicles (Wigge and McMahon, 1998; Slepnev and De Camilli, 2000). Surprisingly, the structure of the amphiphysin BAR domain was found to be highly similar to the structure of arfaptin2 (Tarricone et al., 2001), a protein that interacts with proteins, the strips were incubated with 1:5000 anti-rabbit antibody (1:5000; Amer- then incubated for 1 h with the anti-PICK1 C-100 antibody (Xia et al., 1999). After incubation with proteins, the strips were washed five times in TBS-T and stained for imaging analysis.

Materials and Methods
cDNA cloning, mutagenesis, and protein purification. Rat PICK1 cDNA (Xia et al., 1999) constructs were subcloned into corresponding expression vectors in frames by restriction enzyme digestion and ligation. To generate BAR domain mutants of PICK1, we synthesized PCR primers containing the desired mutation. PCR mutagenesis was performed using a Quickchange site-directed mutagenesis kit (Stratagene, La Jolla, CA). All mutations were subsequently confirmed by sequencing. To produce fusion proteins, cDNA constructs were transformed into Escherichia coli BL21 cells and induced with isopropyl-β-D-thiogalactosidase. Glu- thione S-transferase (GST) fusion proteins were affinity purified by glutathione Sepharose-4B (Amersham Biosciences, Uppsala, Sweden), and maltose-binding protein (MBP) fusion protein was affinity purified by amylose resin (New England Biolabs, Beverly, MA), according to the instructions of the manufacturer. Purified fusion proteins were eluted and dialyzed against the corresponding buffers for follow-up experiments. Fusion protein concentrations were determined by Coomassie assays (Pierce, Rockford, IL).

Lipid binding assay. The lipid binding assay was performed by following the procedure described by Peter et al. (2004). Specifically, brain lipid extracts (Folch fraction B1, B502; Sigma, St. Louis, MO) were resuspended at 2 mg/ml in a buffer containing 20 mM HEPES, pH 7.4, 150 mM NaCl, and 1 mM DTT. The fusion protein (5 μM) was incubated with 0.6 mg/ml liposomes in 100 μl of buffer for 15 min at 37°C and then spun at 140,000 × g for 15 min at 4°C in a Beckman TLA100.1 rotor (Beckman Instruments, Fullerton, CA). After removing all supernatants, the pellets were washed once with the same buffer solution and brought to the same volume as that of the supernatant. The supernatant and the pellet proteins were subjected to SDS-PAGE and visualized by Coomassie stain assays.

Lipid strip assay. The phosphotidylinositol phosphate (PIP) strips (P-6001; Echelon Biosciences, Salt Lake City, UT) were blocked in 3% fatty acid-free BSA in TBS-T (10 mM Tris, pH 8.0, 150 mM NaCl, 0.1% Tween 20) at room temperature for 1 h. The strips were then incubated with 1 μg/ml MBP-PICK1 fusion proteins at room temperature for 2 h. After incubation with proteins, the strips were washed five times in TBS-T and then incubated for 1 h with the anti-PICK1 C-100 antibody (Xia et al., 1999) at room temperature. After washing, the strips were incubated with horseradish peroxidase-conjugated anti-rabbit antibody (1:5000; Amer- sham Biosciences, Buckinghamshire, UK) for 1 h at room temperature, followed by washing in TBS-T. The signals were detected by enhanced chemiluminescence (PerkinElmer Life Sciences, Boston, MA).

Human embryonic kidney 293T cell culture, transfection, and immuno- staining. Human embryonic kidney 293T (HEK293T) cells were cultured in MEM media (Invitrogen, Grand Island, NY) plus fetal bovine serum. For immunostaining, HEK293T cells were grown on coverslips coated with 0.2% gelatin. cDNA constructs of GluR2, green fluorescent protein (GFP)-tagged wild-type PICK1, or mutant PICK1 were transfected into the HEK293T cells by calcium phosphate coprecipitation. The cells were fixed 36–48 h after transfection by 4% paraformaldehyde and 4% sucrose in PBS for 20 min at room temperature. The cells were then permeabilized by 0.2% Triton X-100 in PBS for 10 min at room temperature. After blocking with 10% normal donkey serum (NDS) in PBS for 1 h, the cells were incubated with affinity-purified rabbit anti-GluR2/3 antibody in 3% NDS for 1 h at room temperature, followed by 1 h of incubation with Red-X-conjugated fluorescence anti-rabbit secondary antibody (Jackson ImmunoResearch, West Grove, PA). After washing with PBS, the coverslips were mounted with Permafluor ( Immunon, Pittsburgh, PA). The cells were observed with a Nikon Eclipse TE2000 (Nikon, Tokyo, Japan) inverted fluorescence microscope under a 60× or a 100× Plan Apochromatic oil lens (numerical aperture (NA), 1.4; Nikon). Pictures were taken by a monochromatic low noise cooled CCD camera (SPOT-R; Diagnostic Instruments, Sterling Heights, MI) controlled by MetaMorph imaging acquisition software (Universal Imaging, West Chester, PA). Images were processed with Adobe Photoshop (Adobe Systems, San Jose, CA) to adjust intensity and contrast, to select the region of interest, and to overlay two images. All images were taken in monochromatic gray scale and artificially colored for presentation.

Yeast two-hybrid assay. To test the interaction between GluR2 and PICK1 or PICK1 self-association, the corresponding DNA constructs were subcloned in frames into pPC86 vectors that contained the GAL4 activation domain or pC97 vectors that contained the GAL4 DNA bind- ing domain (Xia et al., 1999). The constructs were cotransformed into H77c yeast cells and grown on double-minus (lacking leucine, trypto- phan) plates. Positive clones were selected and tested on triple-minus plates (lacking leucine, tryptophan, and histidine). After the growth selec- tion, the clones were further tested for β-galactosidase activity.

Coimmunoprecipitation. HEK293T cells were cotransfected with gluta- mate receptor-interacting protein 1 (GRIP1) and myc-PICK1 or myc- PICK1 2-E. Two days after transfection, the HEK293T cells were lysed by 2% Triton X-100 in PBS and incubated with antibody/protein A complex at 4°C for at least 2 h. The resin was washed once with cold PBS and 1% Triton X-100, twice with cold PBS, 1% Triton X-100, and 500 mM NaCl, and three times with cold PBS. After washing, the resin was eluted with 1× SDS sample buffer and was analyzed by SDS-PAGE and immu- noblotted with affinity-purified rabbit anti-GRIP1 or PICK1 antibody.

Generation of Sindbis viruses. The cDNA encoding GFP-tagged wild-type and mutant PICK1 proteins were subcloned into the pSInRep5 viral vector (Invitrogen, Carlsbad, CA). The pSInRep5 constructs and DH26S helper DNA (Invitrogen) were then subjected to restriction enzyme liner- zarization, DNA purification, and in vitro transcription using an SP6 promoter in vitro transcription kit (Ambion, Austin, TX). For production of viruses, mRNAs were transfected into baby hamster kidney cells by Lipofectamine 2000 (Invitrogen). Culturing media were removed 36–48 h after transfection and centrifuged at 2000 g for 10 min at 4°C to remove all cell debris. The supernatants were further centrifuged at 20,000 × g at 4°C for 4 h to harvest the virus particles. The viruses were aliquoted and stored at −80°C. To infect the neurons, the viruses were directly added to the culture media at a titration determined for each batch of viruses. One day after infection, the neurons were fixed and stained for imaging analysis.

Neuronal culture, staining, and quantification. Cultured hippocampal or cortical neurons were prepared from embryonic day 18 Sprague Daw- ley rats and grown on coverslips coated with poly-1-lysine (Sigma). The hippocampal neurons were infected with Sindbis viruses between 16 and 24 d, and the cortical neurons were transfected by calcium phosphate coprecipitation between 13 and 15 d in vitro. For staining, neurons were fixed by 4% paraformaldehyde plus 4% sucrose in PBS for 15 min at 4°C. The neurons were further treated with −20°C methanol for 10 min, washed three times with 0.03% Triton X-100, and then permeabilized by 0.2% Triton X-100 for 10 min at 4°C. After blocking with 10% normal donkey serum for >2 h at room temperature, the neurons were incu- bated with primary antibody in 3% NDS at 4°C overnight to stain their endogenous proteins or 1 h at room temperature for overexpressed protein staining. For endogenous GluR1 and GluR2 staining, antibodies
against the C termini of GluR1 and GluR2 were used, and the GluR2 antibody was directly coupled to Cy3 (Liao et al., 1999). GFP-GluR2-transfected neurons were first incubated with GFP antibody in a medium at 10°C for 30 min to stain the surface GluR2 before fixation. Fluorescent secondary antibodies were added at room temperature for 1 h. After washing, coverslips were mounted and observed under a fluorescence microscope. Quantitative immunofluorescence was performed under a 60× Plan Apochromatic oil lens (1.4 NA; Nikon Instech). Data were acquired and quantified using MetaMorph acquisition and analysis software (Universal Imaging, West Chester, PA). Synaptic clusters were determined by a threshold set at twice the average dendritic gray value. To obtain the surface-to-total GFP-GluR2 ratio, the surface and total gray values of individual clusters were first normalized to dendritic gray to obtain the relative gray value of each individual cluster. The surface-to-total GFP-GluR2 ratio was then obtained by dividing the relative surface gray value of each cluster by its corresponding relative total gray value.

**Biotinylation assay.** HEK293T or COS7 cells were washed three times with PBS supplemented with 0.5 mM CaCl₂ and 0.5 mM MgCl₂ (B buffer) and treated with 0.5 mg/ml sulfo-succinimidyl 6-(biotinamido) hexanoate (sulfo-NHS-LC-biotin; Pierce) in B buffer for 5 min at room temperature or 30 min at 4°C. The free sulfo-NHS-LC-biotin was removed by rapidly washing the cells two times with 100 mM glycine in B buffer followed by two washes with B buffer. The biotinylated cells were solubilized with 1 ml of radioimmunoprecipitation assay buffer (10 mM Tris, pH 7.4, 150 mM NaCl, 1 mM EDTA, 0.1% SDS, 1% Triton X-100, 1% sodium deoxycholate). The samples were centrifuged at maximum speed for 1 h at 4°C with constant rotation. After several washes, the biotinylated surface proteins were eluted from the neutravidin beads in 100 μl of 1× SDS sample buffer. The samples were subjected to SDS-PAGE and Western blot analysis. Instead of using ECL and x-ray film for the Western blot analysis and quantification, we used an Opti-4CN Substrate kit (Bio-Rad, Hercules, CA) to detect protein expression on the polyvinylidene difluoride (PVDF) membrane. The PVDF membrane was incubated in a solution containing 2% Opti-4CN substrate for 15 min and washed several times. In our tests, this method provided better linearity for densitometry analysis compared with the ECL and x-ray film exposure method. The surface GluR2 ratio was calculated only when a linear standard curve was obtained.

**Electrophysiology.** Hippocampal slices were prepared as described previously (Yang et al., 2003). The use and care of animals in this study followed the guidelines of the Shanghai Institutes for Biological Sciences Animal Research Advisory Committee. In brief, postnatal rats (14–17 d of age) were anesthetized with sodium pentobarbital. After decapitation, the hippocampal formations were dissected rapidly and placed in an ice-cold oxygenated (95% O₂ and 5% CO₂) solution containing the following (in mM): 119 NaCl, 2.5 KCl, 1.3 MgSO₄, 1 NaH₂PO₄, 26.2 NaHCO₃, and 11 glucose, pH 7.3. Transverse slices (400 μm thick) were cut with a vibratome (Campden Instruments, Loughborough, UK) and maintained in an incubation chamber for ≥2 h at 25°C before recording. During experiments, individual slices were transferred to a submersion-recording chamber and were continuously perfused with the above-oxygenated solution (3.0 ml/min) at 28°C. Slices were visualized with infrared optics using an Olympus microscope (BX50WI; Olympus Optical, Hamburg, Germany) equipped with DIC optics.

Whole-cell patch recordings were made from CA1 pyramidal neurons in the presence of GABA_A antagonist bicuculline (10 μM). Recording pipettes were filled with a solution containing the following (in mM): 125 Cs-gluconate, 5 CsCl, 10 HEPES, 8 NaCl, 3 Na₂ATP, 0.3 Na-GTP, 0.2 EGTA, 10 Na₂-phosphocreatine, 10 tetraethylammonium-Cl, 5 N-ethyl bromide quaternary salt, pH 7.3. The membrane potential was maintained at −65 to −70 mV. Constant current pulses (10–20 pA, 100 μs, 0.0167 Hz) were applied through extracellular stimulating electrodes (MCE-100; Rhodes Medical Instruments, Woodland Hills, CA) placed at Schaffer collateral of the CA1 region to induce EPSCs in the pyramidal neurons. MBP, wild-type PICK1, and PICK1 2K-E fusion proteins were added into the intracellular solution at a final concentration of 50 μg/ml.

In the experiments to test LTD, data were collected for at least 10 min after breaking-in to allow a full exchange between the cytoplasm and the pipette solutions. LTD was induced with 300 stimuli (1 Hz) at −40 mV (Dudek and Bear, 1992; Mulkey and Malenka, 1992; Daw et al., 2000), with the same intensity as the test stimulus for evoking EPSCs. Data were accepted for analysis only when the postsynaptic currents or voltages and the input resistance did not vary >15% of the average values during the control period. Signals filtered at 5 kHz using the amplifier circuitry were sampled at 10 kHz and analyzed using Clampex 8.2 (Molecular Devices, Union City, CA). Data were presented as means ± SEM, and statistical differences were determined by the Student’s t test. Reagents were obtained from Sigma-Aldrich except where noted.

**Results**

The BAR domain of PICK1 binds to lipids

Because PICK1 is predicted to contain a BAR domain, we first set out to test whether PICK1 binds to lipids. GST-tagged PICK1 fusion protein (ST-PICK1) was incubated with liposomes and then subjected to high-speed centrifugation. Proteins that bind to lipids will be spun down together with liposomes and found in the pellet. We found that GST–PICK1 readily associates with liposomes, because substantial amounts of GST–PICK1 were found in the liposome-containing fraction (top line), but MBP alone was not (bottom line). The lipid binding capability of the BAR domain mutations and the PDZ domain mutant were tested. BAR domain mutants 2K-E, 3K-E, and 5K-E all significantly reduced the binding of PICK1 with lipids, whereas PDZ mutant K270D8-AA did not.

![Figure 1](image-url)

In Figure 1, the BAR domain of PICK1 binds to lipid. A, GST-PICK1s were mixed with brain lipid extract and subjected to high-speed centrifugation to spin down the liposomes. Pellets were redissolved in the same buffer and brought to the same volume as the supernatant. Equal amounts of pellet and supernatant were loaded and resolved by SDS-PAGE. The proteins were then stained by Coomassie blue. A substantial amount of PICK1 was found to accompany the liposome and show up in the pellet (top, left two lanes; S, supernatant; P, pellet.). Without lipid, PICK1 was not found in the pellet (top, right two lanes). As a control, GST itself was incubated with liposome and subjected to the same centrifugation. No GST was found in the liposome-containing pellet (bottom). B, Similar experiments were performed as in A using MBP-PICK1s. Similar to GST-PICK1, MBP-PICK1 was found in the liposome-containing fraction (top line), but MBP alone was not (bottom line). C, The lipid binding capability of the BAR domain mutations and the PDZ domain mutant were tested. BAR domain mutants 2K-E, 3K-E, and 5K-E all significantly reduced the binding of PICK1 with lipids, whereas PDZ mutant K270D8-AA did not.

We then determined the residues of PICK1 that are critical to its lipid binding. The amphiplysin BAR domain forms a
The negative and positive regulation of the lipid binding capability of PICK1

The finding that the 5K-E mutation completely eliminated the lipid binding capability of PICK1 indicates that the BAR domain of PICK1 is necessary for its lipid binding. To test whether the BAR domain of PICK1 is sufficient for lipid binding, we generated the BAR domain only (PICK1–147-358) PICK1 fusion protein and tested its lipid binding capability. We found that the BAR domain itself is fully capable of binding to lipids with slightly higher lipid binding capability compared with full-length PICK1 (Fig. 3).

crescent-shaped dimer, and each monomer uses two pairs of positively charged residues to bind negatively charged membrane (Peter et al., 2004). One pair is located at the end of the crescent dimer, whereas the other pair is located at the concave surface of the dimer. Based on sequence alignments, lysines 266 and 268 of PICK1 are likely to be the pair of positively charged residues located at the end of the crescent dimer and responsible for lipid binding (supplemental Fig. S1 B, available at www.jneurosci.org as supplemental material). We changed both lysines 266 and 268 to negatively charged glutamate and tested the lipid binding capability of the mutant (PICK1 2K-E). As shown in Figure 1C, mutation of lysines 266 and 268 to glutamate eliminated the majority of the lipid binding capability of PICK1. The region of PICK1 corresponding to the second pair of positively charged residues of the amphiphysin BAR domain is more divergent. There are three lysines (K251, 252, 257) that are likely to be responsible for lipid binding (supplemental Fig. S1 B, available at www.jneurosci.org as supplemental material). We substituted glutamate for lysines 251, 252, and 257 and tested the lipid binding ability of the mutant (PICK1 3K-E). As shown in Figure 1C, substitution of glutamate for lysines 251, 252, and 257 also significantly reduced the lipid binding capability of PICK1. We also mutated all five lysines (PICK1 5K-E). As expected, this penta-lysine mutant of PICK1 had minimal lipid binding capability. As a control, we also tested the lipid binding capability of the PICK1 PDZ mutant, in which two critical residues (K27 and D28) of the PDZ domain were changed to alanine. The PICK1 PDZ mutant was found to have similar lipid binding capability to that of wild-type PICK1.

To determine what kinds of lipids interact with PICK1, we performed a lipid strip assay. Purified PICK1 fusion protein was incubated with a lipid strip containing 15 different types of lipid molecules spotted on a nitrocellulose membrane. After washing, PICK1 was detected by immunoblotting. We found that PICK1 preferably binds to phosphoinositides (PtdIns) (Fig. 2A). Specifically, PICK1 binds strongly to PtdIns(3)P, PtdIns(4)P, and PtdIns(5)P. This result was confirmed by probing PIP arrays, which contain a different concentration of phosphoinositides spotted on nitrocellulose membranes. The results from the PIP arrays showed that PICK1 also binds to PtdIns(3)P, PtdIns(4)P, and PtdIns(5)P with relatively high affinity (Fig. 2B). All of the lipid molecules that bind with PICK1 are negatively charged, in agreement with the idea that positively charged residues of the BAR domain interact with negatively charged lipids.
In view of the evidence that the BAR domain only PICK1 binds to lipid better than full-length PICK1, we suspected that other domains of PICK1 may regulate the lipid binding of its BAR domain. The C terminus of PICK1 contains a stretch of negatively charged acidic residues (Fig. 3). One possibility is that this negatively charged acidic domain of PICK1 could bind to the positively charged BAR domain residues and inhibit the lipid binding of the protein. To test this hypothesis, we generated a PICK1 fusion protein with its C-terminal domain deleted (PICK1–1–358). We found that the binding of the PICK1–1–358 fusion protein to lipids was significantly stronger than that of full-length PICK1, indicating that the C terminus of PICK1 indeed serves as an autoinhibitory domain to the lipid binding of PICK1.

To further examine the intramolecular regulation of the lipid binding capability of PICK1, we generated more deletion mutants of PICK1. We found that deleting the PDZ domain of PICK1 (PICK1–110–416) decreased its lipid binding (Fig. 3). We further deleted the 37 amino acid linker region between the PDZ domain and the BAR domain (PICK1–147–416). Deletion of this linker region almost completely diminished the lipid binding of PICK1 in the presence of the C-terminal autoinhibitory domain. It is not clear how the PDZ and the linker region enhance the PICK1 lipid binding capability. The fact that the 5K-E BAR domain mutant does not bind to lipids argues against the possibility that the PDZ domain or the linker region may directly bind to lipids. It is more likely that the PDZ domain and the linker region may affect the conformation of the BAR domain and consequently enhance its lipid binding capability. It has been reported that in certain BAR domains (e.g., the N-BAR domain), a short sequence N terminal to these BAR domains enhances their lipid binding capabilities (Peter et al., 2004). Our result suggests that the BAR domain of PICK1 could be similar to the N-BAR domains found in other proteins.

Lipid binding is critical for subcellular localization and synaptic targeting of PICK1

We then asked what might be the function of PICK1 lipid binding. One possible function is that the lipid binding of PICK1 may regulate its subcellular localization. To test that, we transfected wild-type and lipid binding-deficient mutants of PICK1 into HEK293T cells and examined their subcellular localization. GFP-tagged wild-type PICK1, when transfected into HEK293T cells, was mainly diffuse in the cytosol, with a small number of clusters in some cells (supplemental Fig. S2A, available at www.jneurosci.org as supplemental material). These clusters were usually distributed between the nuclear membrane and the plasma membrane. In contrast, when we transfected lipid binding-deficient mutants into HEK293T cells, these clusters essentially disappeared (supplemental Fig. S2A, available at www.jneurosci.org as supplemental material). We quantified the result and found that the percentages of cells that had at least one cluster were significantly lower for lipid binding-deficient mutants of PICK1 (wild-type PICK1, 40.0 ± 0.3%; 2K-E, 9.2 ± 1.6%; 3K-E, 8.7 ± 1.7%; 5K-E, 9.4 ± 1.2%; p < 0.01) (supplemental Fig. S2B, available at www.jneurosci.org as supplemental material). Similar distribution patterns were also observed when we transfected HEK293T cells with myc-tagged wild-type PICK1 and lipid binding-deficient mutants (data not shown).

In neurons, PICK1 has been shown to be highly enriched at
excitatory synapses (Xia et al., 1999). We wondered whether the lipid binding of PICK1 affects its synaptic targeting. We examined the synaptic localization of lipid binding-deficient mutants of PICK1 compared with wild-type PICK1. GFP-tagged wild-type and lipid binding-deficient mutants of PICK1 were introduced into cultured hippocampal neurons using the Sindbis viral expression system. As shown in Figure 4A, wild-type GFP-PICK1 was highly clustered along the dendrites, mimicking the synaptic localization of endogenous PICK1, and colocalized with PSD-95 at synapses (Fig. 5A). In contrast, the lipid binding-deficient mutants were found to form far fewer clusters in infected neurons (Fig. 4A). We quantified both the number and intensity of PICK1 clusters. The number of clusters of lipid binding-deficient mutants was significantly reduced compared with that of wild-type PICK1 (the number of clusters per 100 μm of dendrite was normalized against wild-type PICK1, which was set at 100%; 2K-E, 37.2 ± 7.5%; 3K-E, 23.6 ± 3.9%; 5K-E, 20.1 ± 3.3%; p < 0.001) (Fig. 4B). The cluster intensities of the 2K-E, 3K-E, and 5K-E mutants were also significantly reduced (the cluster intensity of wild-type PICK1 was normalized to 100%; 2K-E, 49.7 ± 6.0%; 3K-E, 37.7 ± 3.7%; 5K-E, 40.7 ± 5.4%; p < 0.001). We also compared the synaptic targeting of the lipid binding-deficient mutants with PICK1 PDZ mutant (PICK1 KD-AA), which has been shown previously to have reduced synaptic targeting (Boudin and Craig, 2001). We found that although the PICK1 PDZ mutant also has a reduced number and reduced intensity of synaptic clusters, its synaptic targeting is significantly better than that of the lipid binding-deficient mutants (60.3 ± 5.2 and 53.4 ± 5.9% of wild-type PICK1 for cluster number and intensity, respectively; p < 0.001). This result indicates that the lipid binding of PICK1 could be a more fundamental determinant of the synaptic targeting of PICK1. It is interesting to note that deletion of the acidic domain of PICK1 has been reported to enhance its synaptic targeting (Boudin and Craig, 2001). In view of our result that deletion of the acidic domain enhances the lipid binding capability of PICK1, we believe this enhancement of PICK1 synaptic targeting is likely to be the consequence of its increased lipid binding capability. These results indicate that synaptic targeting of PICK1 correlates well with its lipid binding capability.

To rule out the possibility that the change in synaptic PICK1 is a consequence of changing the number of spines or postsynaptic structures, we stained PSD-95, a well-characterized postsynaptic marker, together with wild-type PICK1 and the PICK1 2K-E mutant. We found that wild-type PICK1 colocalized very well with PSD-95 clusters, whereas PICK1 2K-E was essentially missing from these clusters (Fig. 5A). Quantification results show that neither wild-type PICK1 nor PICK1 2K-E mutant significantly changed the number or intensity of PSD-95 clusters (with GFP-infected neurons normalized to 100%, the cluster number per 100 μm of wild-type PICK1, 95.7 ± 3.5%, 2K-E, 93.5 ± 4.6%; the cluster intensity of wild-type PICK1, 93.5 ± 7.1%, 2K-E, 95.6 ± 4.2%; n = 37) (Fig. 5B).

Lipid binding of PICK1 is important for clustering and synaptic targeting of AMPA receptors

We then investigated the function of PICK1 lipid binding in AMPA receptor trafficking. PICK1 has been shown to cluster AMPA receptors when cotransfected into HEK293T cells (Dev et al., 1999; Xia et al., 1999). We wondered whether the lipid binding capability of PICK1 might play a role in this process. As shown in supplemental Figure S3A (available at www.jneurosci.org as supplemental material), when wild-type PICK1 was cotransfected into HEK293T cells together with AMPA receptor subunit GluR2, PICK1 and GluR2 formed many coclusters in discrete sites of the cytosol. The localization of these clusters is similar to that of the PICK1 self-clusters but with greatly increased number and size. The number of these clusters can range from a few to >100 in a single cell. In contrast, when PICK1 2K-E...
was cotransfected with GluR2, we did not observe any coclusters (supplemental Fig. S3B, available at www.jneurosci.org as supplemental material). Similarly, no cocluster with GluR2 was observed for PICK1 3K-E and 5K-E mutants. The quantification result showed that 78.1 ± 4.5% of cells cotransfected with GluR2 and wild-type PICK1 had coclusters, whereas the lipid binding-deficient mutants had no coclusters at all. To rule out the possibility that the loss of GluR2 clustering with PICK1 2K-E was the consequence of lost interaction of PICK1 with GluR2 or loss of PICK1 self-association capability, we performed yeast two-hybrid experiments and found that PICK1 2K-E can still interact with GluR2 and self-associates (supplemental Fig. S4, available at www.jneurosci.org as supplemental material). These results indicate that the lipid binding capability is required for PICK1-induced clustering of AMPA receptors.

In neurons, PICK1 colocalizes with AMPA receptors at excitatory synapses (Xia et al., 1999). Our findings that the lipid binding of PICK1 is required for its synaptic targeting and its clustering of AMPA receptors suggests that lipid binding of PICK1 may also regulate targeting of AMPA receptors to synapses. To test this possibility, we introduced wild-type and lipid binding-deficient mutant PICK1 2K-E into cultured neurons. We found that wild-type PICK1 can increase the synaptic targeting of GluR2 (Fig. 6A). Quantification results showed that PICK1 significantly increased the number and intensity of GluR2 clusters to 125.5 ± 6.7 and 118.3 ± 4.7%, respectively, compared with GFP-infected neurons, which were normalized to 100% (n = 70; p < 0.0) (Fig. 6C). In contrast, the synaptic clusters of GluR2 in neurons infected with lipid binding-deficient mutant PICK1 2K-E were similar to those of GFP-infected neurons but were significantly fewer and weaker compared with those of neurons infected with wild-type PICK1 2K-E (cluster/100 μm, 99.5 ± 6.1%; intensity, 103.1 ± 4.2%; n = 70; p < 0.05 compared with wild-type PICK1) (Fig. 6C).

We also examined the effect of wild-type PICK1 and lipid binding-deficient mutants of PICK1 in synaptic targeting of GluR1, which can form complexes with GluR2. We found that, similarly to GluR2, wild-type PICK1 also significantly increased the number and intensity of synaptic GluR1, whereas the lipid binding-deficient PICK1 2K-E mutant did not (Fig. 6B, D) (the value from GFP-infected neurons was normalized to 100%; GluR1 clusters number per 100 μm, GFP-PICK1, 188.4 ± 12.9%, GFP-PICK1 2K-E, 119.7 ± 9.9%; GluR1 cluster intensity, GFP-PICK1, 153.3 ± 6.2%, GFP-PICK1 2K-E, 97.3 ± 4.8%; n = 30). These results indicate that PICK1 has the capability to target AMPA receptors to synapses, and this capability depends on the lipid binding of PICK1.

**Lipid binding of PICK1 regulates the surface expression of AMPA receptors**

PICK1 has been found to regulate the surface expression of a number of proteins. We tested whether the lipid binding of PICK1 was involved in this process. This was done by transfecting wild-type PICK1 or PICK1 2K-E with GluR2 into HEK293T cells. Surface GluR2 was labeled with membrane-impermeable biotin. After lysing the cells, surface GluR2 was separated from intracellular GluR2 via binding of biotin to immobilized neutravidin. Above, the intensity and number of GluR2 clusters per 100 μm of dendrite length were quantified from multiple experiments. The number and intensity of GluR2 clusters from GFP-PICK1-infected neurons were significantly higher than neurons infected with GFP only or PICK1 lipid binding-deficient mutant 2K-E (p < 0.001). D, The intensity and number of GluR1 clusters were quantified from neurons infected with GFP, GFP-PICK1, and GFP-PICK1 2K-E. Wild-type PICK1 overexpression significantly increased the GluR1 cluster number (***p < 0.001), but the lipid binding-deficient mutant 2K-E did not ( p > 0.05). Error bars represent SEM.
Reduced the surface ratio of GluR2 (GluR2 alone, 14.6 ± 0.4%; GluR2 plus PICK1-wild type, 9.4 ± 0.1%; n = 6; p < 0.01) (Fig. 7B). In contrast, PICK1 2K-E mutant did not significantly alter the surface expression of GluR2 (PICK1–2K-E, 14.9 ± 1.3%; n = 6; p > 0.05). These results indicate that lipid binding is important for PICK1-regulated surface expression of AMPA receptors.

To further investigate the role of PICK1 lipid binding in surface expression of AMPA receptors at synapses, we performed surface staining of GluR2 and measured the surface-to-total ratio of GluR2 at the synapses. GFP-tagged GluR2 was transfected into cortical neurons together with myc-tagged wild-type PICK1 or PICK1 2K-E mutant. Surface GluR2 was labeled with GFP antibody, which was added to the neurons before fixation. After fixation, signal from the GFP antibody was visualized as surface GluR2, whereas the native GFP signal was taken as total GluR2 (Fig. 7C). The intensities of both surface and total GluR2 clusters were measured and normalized to respective dendritic gray values to obtain the relative intensities. The surface-to-total ratio of GluR2 clusters was determined by dividing the relative surface intensity by the total intensity of individual clusters. We found that wild-type PICK1 significantly decreased the ratio of GluR2 on the synaptic surface, whereas PICK1 2K-E mutant did not (surface:total ratio: vector control, 2.36 ± 0.02; wild-type PICK1, 2.19 ± 0.04; PICK1 2K-E, 2.36 ± 0.02; n = 30) (Fig. 7D). The findings that PICK1 increases the total (Fig. 6) but reduces the surface ratio of AMPA receptors at synapses suggest that PICK1 maintains an intracellular reserve pool of AMPA receptors at synapses in a lipid binding-dependent manner.

Figure 7. Lipid binding of PICK1 is important for the surface expression of GluR2. A, Different cDNA constructs were transfected into HEK293T cells as indicated. Surface GluR2 was isolated using the surface biotinylation assay. Total and surface samples were resolved by SDS-PAGE and immunoblotted with anti-GluR2/3C antibody. As indicated by the intensity of surface GluR2 band, wild-type PICK1 notably reduced the surface expression of GluR2, whereas PICK1–2K-E did not. B, Quantification data from multiple experiments. Surface GluR2 ratios, which are expressed as the percentage of GluR2 on the surface relative to the total GluR2, were determined by fitting the surface GluR2 to a standard curve obtained by loading different amounts of total GluR2 (**p < 0.01). C, GFP-GluR2 was transfected into neurons together with the vector control, wild-type PICK1, or PICK1 2K-E mutant. The surface GluR2 (red) was stained with GFP antibody added to neurons before fixation. The total GluR2 (green) was obtained from native GFP signal. Scale bar, 10 μm. D, Quantification data from multiple experiments. The surface-to-total ratio of GluR2 at the synapses was significantly lower in wild-type PICK1 transfected neurons compared with neurons infected with the vector control or PICK1 2K-E mutant (*p < 0.05). E, A model illustrates that PICK1 (green) maintains an intracellular reserve pool of AMPA receptors at the synapses by binding to negatively charged lipids (red color). Overexpression of PICK1 increases the intracellular synaptic pool by internalizing GluR2 containing AMPA receptors (pink) from synaptic surface. Extrasynaptic AMPA receptors, especially GluR1 homomeric receptors (yellow), then move laterally to the synaptic surface to replace the internalized receptors. The end results of PICK1 overexpression are to increase the total AMPA receptors at the synapses. Although both GluR1 and GluR2 are increased in synapses, their distribution at the surface, however, could be differentially affected by PICK1. Although PICK1 reduces surface synaptic GluR2, it may not change or even increase surface synaptic GluR1 with the addition of homomeric GluR1 receptors. Error bars represent SEM.
The results from experiments shown in Figure 8. Lipid binding of PICK1 is important for long-term depression. A, Application of the PICK1 2K-E mutant (open circles) through patch pipettes increased the EPSC amplitude that reached a steady-state level within 10 min. Data were collected immediately after the establishment of whole-cell recording and normalized in each neuron by the mean amplitude of EPSCs observed during the initial 2 min of recording. B, Summary of the results from the experiments shown in A. Data represent averaged EPSCs recorded during the period from 7 to 15 min after the establishment of whole-cell recording and are normalized as in A. The number associated with each column refers to the number of neurons tested in each condition. C, Inhibition of low-frequency stimulation (LFS)-induced LTD by the PICK1 2K-E mutant. EPSCs were recorded 10 min after the establishment of whole-cell recording and normalized by the mean amplitude of EPSCs recorded during the control period (before LFS application). D, Summary of the results from experiments shown in C. Data represent averaged EPSCs recorded 20 min after LFS application and are normalized as in C. The number associated with each column refers to the number of neurons tested in each condition. A, C, Insets, Sampled traces of averaged EPSCs in the PICK1 2K-E group (A) and the control group (C), respectively, recorded during the period indicated by the corresponding numbers. Calibration: A, 50 pA, 20 ms; C, 50 pA, 30 ms. **p < 0.01; ***p < 0.001. Error bars represent SEM.

Discussion

In this study, we demonstrated that PICK1 can directly bind to lipids via its BAR domain. The lipid molecules that bind to PICK1 are mainly negatively charged phosphoinositides. Mutations of critical residues within the PICK1 BAR domain disrupt its lipid binding. Moreover, lipid binding of the BAR domain is negatively regulated by the acidic domain and positively regulated by the PDZ domain and the linker region of PICK1. We also provide evidence for two potential physiological functions of PICK1 lipid binding. The first is that lipid binding is critical to the subcellular localization of PICK1 in cells. In HEK293T cells, lipid binding-deficient mutants of PICK1 do not concentrate in discrete cytosolic sites as wild-type PICK1 does. As a consequence, these mutants cannot cluster the PICK1 PDZ binding partner GluR2 in HEK293T cells. In neurons, the lipid binding-deficient mutants do not target to synapses as wild-type PICK1 does. We found that whereas PICK1 can increase synaptic targeting of AMPA receptors, lipid binding-deficient PICK1 loses the ability to target AMPA receptors to synapses. The second function is that PICK1 lipid binding is important for the surface expression of AMPA receptors. Without the lipid binding capability, PICK1 loses its ability to regulate surface expression of AMPA receptors, which leads to deficiency in long-term depression. This suggests a mechanism for PICK1-mediated AMPA receptor trafficking: although PICK1 PDZ domain binds to the C termini of AMPA receptors, its BAR domain binds to negatively charged lipids and tethers the AMPA receptors as cargo to curved-membrane microdomains that are designated to form trafficking vesicles (Fig. 7E). We believe this mechanism may not be limited to PICK1-mediated AMPA receptor trafficking. Because PICK1 interacts with a number of membrane proteins, it might be a general mechanism for PICK1 to regulate the trafficking of membrane proteins via the lipid binding capability of its BAR domain.

It has been reported that overexpression of PICK1 leads to reduction of surface GluR2 at synapses (Perez et al., 2001; Terashima et al., 2004). Our result that PICK1 increases the total but reduces the surface-to-total ratio of AMPA receptors at synapses in a lipid binding-dependent manner suggests that PICK1 maintains an intracellular pool of AMPA receptors at synapses by binding to negatively charged lipids (Fig. 7E). This intracellular pool of AMPA receptors maintained by PICK1 could be the reservoir for AMPA receptor traffic to and from synaptic surface during synaptic plasticity (Daw et al., 2000; Xia et al., 2000). This notion is supported by the data that disruption of the PICK1 interaction with GluR2 or the PICK1 lipid binding capability impairs LTD, because these manipulations block the movement of AMPA receptors from the surface to the intracellular pool at synapses. The model is also in agreement with the recent finding that PICK1 knock-out mice are deficient in the activity-dependent insertion of GluR2 containing AMPA receptors (Gardner et al., 2005), which could be explained by a depletion of the intracellular reserve pool of AMPA receptors in PICK1 knock-out mice. It is interesting to note that both synaptic targeting and surface expression of AMPA receptors are dependent on the lipid binding of PICK1. It is conceivable that during LTP or LTD induction, certain intracellular signal transduction pathways may change the lipid composition or affinity of PICK1 to thus regulate the ratio between the surface and intracellular pool of AMPA receptors at synapses. In fact, one lipid kinase, phosphatidyl inositol 3-kinase, has been reported to be involved in AMPA receptor trafficking during synaptic plasticity (Man et al., 2003).

In this study, we provided the first evidence that PICK1 increases both total GluR2 and GluR1 at synapses (Fig. 6). Although GluR1 does not directly interact with PICK1, the majority of GluR1 in neurons is coupled with GluR2 and is likely to traffic together with GluR2. However, this finding seems at odds with previous reports that PICK1 only affects surface GluR2, not GluR1 (Terashima et al., 2004; Lu and Ziff, 2005). It is even more surprising that PICK1 increases total GluR1 more than GluR2. However, this seeming discrepancy can be explained by the model that PICK1 maintains an intracellular synaptic pool of AMPA receptors, and this pool exchanges AMPA receptors with the surface synaptic pool and surface extrasynaptic pool. When overexpressing PICK1, excessive PICK1 will move GluR1 and
GluR2 heteromeric receptors from the surface synaptic pool to the intracellular synaptic pool. Extrasynaptic AMPA receptors, especially GluR1 homomeric receptors, then move to the synaptic surface by lateral diffusion to replace the internalized receptors. This leads to an increase of total AMPA receptors, especially GluR1, at the synapses and decrease of surface GluR2 but not surface GluR1 at the synapses (Fig. 7E). This model is also in agreement with that reported by Tereshima et al. (2004) in which overexpression of PICK1 reduced surface GluR2 but increased synaptic AMPA current and inward rectification, indicating more GluR1 homomeric receptors moved to the synaptic surface. However, it should be noted that the larger effect of PICK1 on GluR1 than GluR2 should be taken with caution. We noticed that the GluR2 antibody has a high background signal on the dendrites and does not stain synapses as strongly as GluR1 antibody. The high background level of GluR2 antibody may have averaged down the signal and made the effect of PICK1 on GluR2 appear to be smaller. Additional studies are required to further clarify the role of PICK1 in GluR1 and GluR2 trafficking.

In a recent study, the BAR domain of PICK1 was found to interact with GRIP/ABP, a PDZ domain-containing protein that also interacts with GluR2/3 (Lu and Ziff, 2005). The interaction of PICK1 and GRIP/ABP was also found to be important for AMPA receptor trafficking. We found that PICK1 2K-E, the lipid binding-deficient mutant used in this study, binds to GRIP/ABP in a similar way as wild-type PICK1 (supplemental Fig. S3, available at www.jneurosci.org as supplemental material). This indicates that the effect we observed using the PICK1 2K-E mutant is likely a result of lipid binding, instead of disrupting the interaction of PICK1 with GRIP/ABP. Apparently, the lipid binding region of the PICK1 BAR domain is distinct from its binding region with GRIP. It would be interesting to determine in future studies whether there was any interplay between PICK1 interaction with lipids and its interaction with GRIP/ABP.

Many of the BAR domain-containing proteins, such as amphiphysin and endophilins, are accessory proteins in endocytosis (Marsh and McMahon, 1999; Slepelev and De Camilli, 2000; Peter et al., 2004). These accessory proteins do not directly bind to cargo proteins during endocytosis. Instead, they form complexes with the core endocytotic proteins. For example, amphiphysin was found to interact with AP2, clathrin, dynamin, and synaptopodin to regulate the endocytosis of synaptic vesicles (Wigge and McMahon, 1998; Slepelev and De Camilli, 2000). The BAR domain of amphiphysin can sense and bend the lipid bilayer to help form trafficking vesicles (Peter et al., 2004). In contrast to amphiphysin, PICK1 has a PDZ domain in addition to its lipid binding BAR domain. The PDZ domain is a well-characterized protein motif that interacts with the C termini of many membrane proteins (Sheng and Sala, 2001). This suggests that PICK1, unlike other BAR domain proteins, could directly bind to cargo proteins for endocytosis.

The interaction with synaptic scaffolding proteins has been proposed as an important mechanism for targeting of synaptic proteins. For example, PSD-95, the prototypical PDZ domain-containing protein and a major component of postsynaptic density, plays a central role in assembling the postsynaptic protein complex by interacting with a number of receptors, signaling molecules, and proteins linked to the cytoskeleton (Sheng and Sala, 2001). Protein–protein interactions have also been found to be important for synaptic targeting of PICK1. Mutations of the PDZ domain and the coiled-coil domain, both are protein–protein interaction motifs, reduce the synaptic clusters of PICK1 (Boudin and Craig, 2001). Our finding that the lipid binding of PICK1 is required for its synaptic targeting provides a new mechanism for targeting proteins to synapses. It should be noted that lipids have been found previously to be important for targeting of synaptic proteins. PSD-95 was found to be palmitoylated and integrated into membranes after palmitoylation (Craven et al., 1999). This palmitoylation was found to be important for synaptic targeting of PSD-95 and regulation of synaptic strength (Topinka and Bredt, 1998; El-Husseini Ael D, et al., 2002). In contrast to the palmitoylation of PSD-95, which covalently links the protein with fatty acid, the interaction of the PICK1 BAR domain with lipids is based on a noncovalent interaction, which makes the association of PICK1 with lipids more dynamic. It is conceivable that PICK1 can be recruited to a specific membrane region by adding negatively charged lipids and also easily dissociated when the membrane becomes less charged. This gives PICK1 the ability to regulate the surface expression of membrane proteins at specific times and locations, which is crucial for physiological functions such as synaptic plasticity.

References


