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Journal Title: Cellular Physiology and Biochemistry
Volume: Volume 30, Number 6
Publisher: Karger: Open Access Journals | 2012, Pages 1444-1455
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1159/000343332
Permanent URL: http://pid.emory.edu/ark:/25593/fk18t

Final published version: http://www.karger.com/Article/Abstract/343332

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Accessed March 5, 2020 3:49 PM EST
PSD-95 Interacts with NBCn1 and Enhances Channel-like Activity without Affecting Na/HCO₃ Cotransport

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Key Words
Bicarbonate transport • Acid-base • pH • Electrophysiology • Neuron • Xenopus

Abstract
Background/Aims: The sodium/bicarbonate transporter NBCn1 plays an essential role in intracellular pH regulation and transepithelial HCO₃⁻ movement in the body. NBCn1 also has sodium channel-like activity uncoupled to Na/HCO₃ cotransport. We previously reported that NBCn1 interacts with the postsynaptic density protein PSD-95 in the brain. Here, we elucidated the structural determinant and functional consequence of NBCn1/PSD-95 interaction.

Methods: Results: In rat hippocampal CA3 neurons, NBCn1 was localized to the postsynaptic membranes of both dendritic shafts and spines and occasionally to the presynaptic membranes. A GST/NBCn1 fusion protein containing the C-terminal 131 amino acids of NBCn1 pulled down PSD-95 from rat brain lysates, whereas GST/NBCn1-ΔETSL (deletion of the last four amino acids) and GST/NBCn2 (NCBE) lacking the same ETSL did not. NBCn1 and PSD-95 were coimmunoprecipitated in HEK 293 cells, and their interaction did not affect the efficacy of PSD-95 to bind to the NMDA receptor NR2A. PSD-95 has negligible effects on intracellular pH changes mediated by NBCn1 in HEK 293 cells and Xenopus oocytes. However, PSD-95 increased an ionic conductance produced by NBCn1 channel-like activity. This increase was abolished by NBCn1-ΔETSL or by the peptide containing the last 15 amino acids of NBCn1.

Conclusion: Our data suggest that PSD-95 interacts with NBCn1 and increases its channel-like activity while negligibly affecting Na/HCO₃ cotransport. The possibility that the channel-like activity occurs via an intermolecular cavity of multimeric NBCn1 proteins is discussed.

Introduction

NBCn1 (SLC4A7) is an ion transporter that normally moves Na⁺ and HCO₃⁻ into cells [1, 2]. NBCn1 contributes to transepithelial movement of ions and acid/base equivalents in epithelial cells and intracellular pH (pHᵢ) regulation in non-epithelial cells [3, 4]. Slc4a7 knockout mice
Lee/Yang/Kim et al.: PSD-95/NBCn1 Interaction and its Functional Effect on NBCn1

Cellular Physiology and Biochemistry

Materials and Methods

All experiments in this study were conducted under the National Institute of Health guidelines for research on animals, and experimental protocols were approved by the Institutional Animal Care and Use Committee at Emory University.

Electron microscopic immunoperoxidase

Electron microscopic immunohistochemistry was performed using the previous protocol [16] with slight modification. Briefly, adult rats (Sprague Dawley from Harlan; Indianapolis, IN, USA) were fixed in 4% paraformaldehyde and 0.1% glutaraldehyde, and brains were cut into 10-mm-thick blocks. Hippocampal sections (60 µm) were treated in 1% sodium borohydride and then placed in a cryoprotectant solution (25% sucrose and 10% glycerol) for 20 min and then frozen at -80°C for 20 min. The sections were thawed and treated with a serial dilution of cryoprotectant (100–30%). After washes in PBS, the sections were treated with 10% normal goat serum and 1% bovine serum albumin, and incubated with 1:200 diluted NBCn1 antibody [17] at 4°C. The sections were incubated with biotinylated goat anti-rabbit IgG (1:200; Vector Laboratories; Burlingame, CA, USA) for 1.5 h and then the DAB staining was done. Post-fixation was done in 1% osmium tetroxide and dehydration in a serial dilution of ethanol and propylene oxide. The sections were embedded in epoxy resin Durcupan ACM (Sigma-Aldrich; St. Louis, MO, USA), incubated at 60°C for 2 days, and glued to the resin blocks. Ultrathin sections were collected onto Pioloform-coated single-slot copper grids and stained with lead citrate. The sections were visualized with a Zeiss EM 10C electron microscope. Images were analyzed DigitalMicrograph software (Gatan; Pleasanton, CA, USA).

Double-label immunofluorescence

Double label of NBCn1 and PSD-95 immunofluorescence was performed as described previously [18]. Mouse brains were fixed with ice-cold 4% paraformaldehyde and brains sections (30 µm) were washed...
with PBS containing 0.1% Tween 20 and then blocked with 10% normal goat serum for 1 h. The sections were incubated with the rabbit NBCn1 antibody (1:100) and the mouse PSD-95 antibody (1:100) at 4°C overnight. The secondary antibodies were Alexa 488 anti-rabbit IgG (1:500; Invitrogen; Carlsbad, CA) and Alexa 594 anti-mouse IgG (1:500; Invitrogen) for 1 h incubation. The sections were mounted onto gelatin-coated slides with Vectashield (Vector Laboratories; Burlingame, CA, USA), covered with coverslips, and visualized using an Olympus Fluoview FV1000 confocal microscope with UPLFLN objective (numerical aperture 1.3). Images of CA3 neurons were acquired using Olympus software.

**Pull-down assay**

The following GST fusion proteins were prepared: GST/NBCn1 containing the C-terminal 131 amino acids of rat NBCn1-E; GST/NBCn1-ΔETSLS that lacks the last 4 amino acids ETSLS of NBCn1-E; GST/NBCn2 containing the C-terminal 113 amino acids of rat NBCn2-C (i.e., rb2NCBE). Fusion proteins were constructed by subcloning the corresponding nucleotides into pGEX-4T (GE Healthcare; Chicago, IL, USA). Brains of adult rats were homogenized in the lysis buffer (300 mM mannitol, 0.1 mg/ml phenylmethanesulphonyl fluoride, 5 mM HEPES, 1 μM protease inhibitor cocktail, pH 7.2) and centrifuged at 3,000 × g for 10 min to remove cell debris. The lysates were then incubated with GST only or GST fusion proteins bound to glutathione-Sepharose beads for 4 h. The beads were washed 4 times with 0.5% Nonidet P-40 in PBS, and proteins bound to the beads were dissociated by adding the SDS-PAGE sample loading buffer. Immunoblot was performed with the mouse PSD-95 antibody (1:500; Affinity Bioreagents; Golden, CO, USA). In parallel experiments, GST only or GST fusion proteins bound to glutathione-Sepharose beads were loaded on the gel and stained with Coomassie blue to visualize the molecular weight of the fusion proteins.

**Immunoprecipitation**

HEK 293 cells were transfected with pcDNA3.1/NBCn1, GW1/myc-PSD-95, pcDNA3.1/NR1A, and pcDNA3.1/NR2A using Lipofectamine 2000 (Life Technologies; Grand Island, NY) according to the manufacturer’s protocol. The amounts of plasmids were 4 μg NR1A/NR2A, 4 μg PSD-95, and 0.5–4 μg NBCn1 to transfect 2–6 × 10^6 cells. NR1A was omitted in some experiments as it did not significantly affect NR2A/PSD-95 interaction. pcDNA3.1 vector served as a control. At 48 h after transfection, cells were lysed in 20 mM HEPES, 150 mM NaCl, 2 mM EDTA, 10% glycerol, 0.5% Nonidet P-40, and 1 × protease inhibitor cocktail, and centrifuged at 13,200 × g for 15 min. Supernatants were pre-treated with Protein A/G agarose beads at 4°C for 20 min to block non-specific bead binding. Beads were removed, and the lysates were incubated with the PSD-95 antibody at 4°C overnight in 100 mM NaCl, 0.5% Triton X-100, 0.2% sodium deoxycholate, and 20 mM HEPES, pH 7.4. Fresh beads were added to the lysates and agitated for 1 h. Beads were collected and washed, and proteins were dissociated from the beads by adding the SDS-PAGE sample loading buffer. Immunoblot was performed with the NBCn1 antibody and the NR2A antibody (Millipore; Billerica, MA, USA).

**Immunoblotting**

HEK 293 cells and oocytes were homogenized with a 26 gauge needle in ice-cold lysis buffer and centrifuged at 810 × g for 10 min to remove cell debris. Crude plasma membranes were prepared by centrifugation at 100,000 × g for 30 min at 4°C. Protein concentration in the membrane preparation was quantitated using the Bradford reagents (Sigma-Aldrich). The equal amounts of protein samples (10–30 μg) were loaded on a 7.5% SDS-PAGE and blotted to a nitrocellulose membrane. The blot was treated with blocking buffer containing 0.05% Tween 20 and 5% non-fat dry milk in PBS for 1 h and then with primary antibodies for 1 h. After washes with PBS containing 0.05% Tween 20 for 5 min, the blot was treated with a horseradish peroxidase-conjugated anti-rabbit or anti-mouse secondary antibody for 1 h (1:2500; Millipore). After washes, the blot was treated with the chemiluminescence detection solution from the ECL kit (GE Healthcare). The blot was stripped and reprobed for β-actin. For densitometric comparison, pixel intensities of the immuno-reactive bands were measured by positioning boxes around the bands using ImageJ image analysis software (NIH; Bethesda, MD, USA). NBCn1 or NR2A were normalized to β-actin after background subtraction.

**Measurements of pH in HEK 293 cells**

Cells growing on a coverslip (2.2 × 10^5 cells in a 60-mm dish) were transfected as described above and incubated for 36 h. Cells were loaded with 6.5 μM of bis-carboxyethyl carboxyfluorescein acetoxymethyl ester (BCECF-AM) for 10 min, and the coverslip was mounted to a chamber RC-30 (Harvard Apparatus; Holliston, MA, USA) affixed on the stage of a Nikon TE200 inverted microscope. Cells were perfused with HEPES-buffered solution (mM: 140 NaCl, 1 KCl, 1 MgSO4, 2 CaCl2, 2.5 NaH2PO4, 5.5 glucose, and 10 HEPES, pH 7.5). The HCO3−/CO2-buffered solution was made by replacing 33 mM NaCl in HEPES-buffered solution with 33 mM NaHCO3, 5% CO2 (pH 7.5). The two solutions contained 10 μM ethyl isopropyl amiloride to inhibit endogenous Na/H exchanger. At the end of the experiments, cells were exposed to a high K+/nigericin solution [19] of pH 6.8. The dye was alternately excited with 500 nm and 440 nm light using a Lambda 10-2 filter wheel controller (Sutter Instruments; Novato, CA, USA) and the 530 nm emission lights from both excitations were captured. The ratio of the emission intensities at 500 nm to 440 nm was calculated and normalized to the ratio in the K+ /nigericin solution. Background subtraction was made from the area without
cells. The ratio was converted to pH using a calibration curve constructed from the ratios at different pH, ranging from 6.0 to 8.0. Data were acquired using Metafluor software (Universal Imaging, Downingtown, PA, USA). The rate of pH changes (i.e., dpH/dt) was calculated from the slope of a linear regression line during the first 30 seconds of recovery. The degree of CO₂-induced acidification was similar among cells expressing NBCn1 only or NBCn1/PSD-95. All experiments were performed at 37°C.

Protein expression in Xenopus oocytes
Oocytes were prepared as described previously [7]. Briefly, frogs were purchased from Xenopus Express (Brooksville, FL, USA). A frog was anesthetized with fresh 0.1% 3-aminobenzoic acid ethyl ester for 20 min and surgery was done below the termination of sternum to collect ovary. After surgery, the frog was returned to a recovery tank containing 0.1 M NaCl. Oocytes were agitated with Ca²⁺-free ND96 solution (mM; 96 NaCl, 2 KCl, 1 MgCl₂, and 10 HEPES, pH 7.5) for 5 times for 20 min each, and then with type IA collagenase (2 mg/ml; Sigma-Aldrich) twice for 20 min each. Stage V-VI oocytes were sorted under the dissection microscope and stored at 18°C overnight. For injection, cRNAs of NBCn1, PSD-95, and NR1A/NR2A were made using the mMessage/mMachine kit (Life Technologies). The amount of injected cRNA (in 46 nl) was 11 ng for NBCn1, 7.5 ng for PSD-95, 5 ng for NR1A and 12.5 ng for NR2A. Thus, the ratio of NR1A to NR2A was 1 to 2.5. For pH measurements, 35 ng for each NBCn1 and PSD-95 were used. Control oocytes were injected with sterile water or uninjected. Oocytes were maintained at 18°C for 3–4 days before use.

Measurements of pH in oocytes
Oocyte pH was measured using the protocol [20] with slight modification. A glass capillary was filled with the proton ionophore 1 cocktail B (Sigma-Aldrich) and back-filled with the phosphate buffer (pH 7.0). This pH electrode was connected to a high-impedance electrometer FD-223 (World Precision Instruments, Sarasota, FL, USA) routed to a custom-made subtraction amplifier. A second glass capillary filled with 3 M KCl was connected to a voltage clamp amplifier OC-725C (Harvard Apparatus). Signals were sampled by a Digidata 1322A (Molecular Devices; Sunnyvale, CA) and data were acquired using pClamp 8 (Molecular Devices). The signal in voltage electrode was subtracted from that in pH electrode to calculate the voltage for pH. The slope of voltage to pH was determined using pH 6.0 and 8.0 standards (typically 53 ± 3 mV/pH).

Two-electrode voltage clamp
Two-electrode voltage clamp was performed to record the steady-state current produced by NBCn1. A glass capillary was filled with 3 M KCl and connected to a voltage headstage, and another capillary was connected to a current cable (tip resistance of 1–2 MΩ for each). The voltage and current cables were then connected to OC-725C. Oocytes were impaled with the two electrodes in ND96 solution and clamped at −60 mV. After the holding currents became stable, voltage commands from −120 mV to +60 mV with 20 mV increments were applied with the duration of 100 msec. Current-voltage (I–V) relationships were then done 36 h later to calculate NBCn1-associated ionic conductance. The one-way ANOVA with Bonferroni post test was used to analyze I) pH, recovery rate and slope conductance of NBCn1 affected by PSD-95 and ii) slope conductance of NBCn1 or NBCn1/PSD-95 affected by the peptide. The p value of less than 0.05 was considered significant.

Results

NBCn1 is predominantly localized in dendritic shafts and spines
We performed electron microscopic immunoperoxidase staining of rat hippocampal CA3 neurons to determine synaptic expression of NBCn1. NBCn1 is localized to both dendrites and cell bodies at the light microscopic level [13, 22]. PSD-95 is predominantly localized to dendritic spines at the light and electron microscopic levels [23]. Thus, our experiments were done with the apical proximal dendrites where glutamatergic synapses are heavily
innervated. Fig. 1A shows electron micrographs of the stratum radiatum in cross section. NBCn1 immunoperoxidase staining was mostly confined to postsynaptic membranes. Both dendritic spines and shafts were positive for NBCn1. In addition, presynaptic membranes were occasionally stained. Fig. 1B shows confocal images of proximal dendrites double-labeled with antibodies to NBCn1 (green; probed with Alexa 488 fluorescent dye) and PSD-95 (red; probed with Alexa 594 fluorescent dye). NBCn1 was prominently dispersed along dendritic processes, whereas PSD-95 was mostly in puncta. The two fluorescence labelings frequently overlapped in large dendrites.

**NBCn1 interacts with PSD-95 via the C-terminal PDZ motif**

We previously demonstrated the interaction of NBCn1/PSD-95 in rat brain lysates by coimmunoprecipitation and GST pull-down assay [13]. To further assess the specificity of the interaction between NBCn1 and PSD-95, we performed pull-down assays in which rat brain lysates were incubated with GST/NBCn1 containing the C-terminal 131 amino acids of rat NBCn1, GST/NBCn2 containing the C-terminal 113 amino acids of rat NBCn2-C (i.e., rb2NCBE) and GST/NBCn1-ΔETSL (the deletion of the last 4 amino acids ETSL in NBCn1). Probed with the PSD-95 antibody, the immunoblotting of pull-down samples detected a protein band (95 kDa) in GST/NBCn1 (Fig. 2A). The antibody recognized an additional weak band of ~75 kDa protein that was also present in the lysates. PSD-95 was not detected in GST alone and GST/NBCn2. GST/NBCn1-ΔETSL was also negative, consistent with the idea that the NBCn1/PSD-95 interaction is mediated by the PDZ motif in the NBCn1 C-terminus. The interaction of NBCn1/PSD-95 was confirmed by coimmunoprecipitation using HEK 293 cells (Fig. 2B). Lysates from cells transfected with the two proteins were immunoprecipitated with the PSD-95 antibody and then immunoblotted with the NBCn1 antibody. The immunoblotting recognized NBCn1 in the precipitates from cells expressing NBCn1/PSD-95. The control cells expressing NBCn1 alone showed no NBCn1 in the precipitates.

Among proteins capable of binding to PSD-95, NMDA receptors are the major protein that constitutes the macromolecular complex with PSD-95 in the postsynaptic density [24, 25]. To test whether NBCn1/PSD-95 interaction affects the efficacy of PSD-95 to interact with NMDA receptors, we expressed NBCn1, PSD-95, and the NMDA receptor subunits NR1A and NR2A (hereafter NR1A + NR2A are referred to as NMDAR) in HEK 293 cells and performed coimmunoprecipitation (Fig. 2C and D). Cell lysates were immunoprecipitated with the PSD-95 antibody and then immunoblotted either with the NBCn1 antibody or the NR2A antibody. We found both NBCn1 and NR2A in the precipitates from cells expressing NBCn1/NMDAR/PSD-95 (Fig. 2C; lane 2). This detection was not due to incomplete wash
of Protein A/G beads after the lysate incubation because the immunoreactive bands were absent from cells expressing NBCn1 and NMDAR, but not PSD-95. Coimmunoprecipitation was then repeated with cells expressing PSD-95, NMDAR, and different amounts of NBCn1 (Fig. 2D). The NBCn1 signal intensity was progressively stronger at the higher amounts of NBCn1 for transfection, whereas the NR2A intensity remained unaffected under the same condition. The similar intensity in all lanes demonstrates that NBCn1/PSD-95 interaction occurs without affecting the PSD-95/NMDAR interaction.

**PSD-95 has negligible effects on NBCn1-mediated pH change**

To examine the functional effect of PSD-95 on NBCn1, we measured pH of HEK 293 cells expressing NBCn1 and PSD-95 or vector using the pH-sensitive fluorescence dye BCECF-AM. NBCn1 moves HCO$_3^-$ into the cells and raises pH, and thus experiments were focused on monitoring pH recovery from an acid load in the presence of HCO$_3^-$/CO$_2$. Solutions contained 100 μM EIPA to block endogenous Na/H exchanger activity. Fig. 3A shows representative pH traces during the application of 33 mM HCO$_3^-$, 5% CO$_2$ (pH 7.5). Applying HCO$_3^-$/CO$_2$ caused an abrupt fall in pH as CO$_2$ entered into cells and produced H$^+$ after hydration. The pH was then recovered from the CO$_2$-induced acidification as NBCn1 moved HCO$_3^-$ into the cells and buffered H$^+$. The pH change during this recovery was similar in the two groups of cells expressing NBCn1/PD-95 and NBCn1 alone. The resting pH and CO$_2$-induced acidification were also similar in these cells, reflecting that the buffering powers are not different. Fig. 3B summarizes the mean rate of pH change (dPH /dt) during the recovery from acidification (n = 6–8 cells from one coverslip for each group; triplicate experiments). No significant change in dPH /dt was found regardless of PSD-95 (110 ± 13 × 10$^{-6}$ pH units/sec for NBCn1 and 114 12 × 10$^{-5}$ pH units/sec for NBCn1/PD-95; p > 0.05).

To further confirm the negligible effect of PSD-95 on NBCn1-mediated pH changes, we expressed NBCn1 and PSD-95 in *Xenopus* oocytes and measured pH using a proton-selective microelectrode. In an oocyte expressing PSD-95 (Fig. 4A), the pH was not recovered from CO$_2$-induced acidification and instead reached steady-state. This confirms that PSD-95 alone does...
not induced pH recovery. In an oocyte expressing NBCn1 (Fig. 4B), the pH falls upon exposure to HCO$_3^-$ /CO$_2^-$, but was then recovered from the acidification as NBCn1 moved HCO$_3^-$ into the oocyte and buffered intracellular H$^+$. In an oocyte coexpressing the two proteins (Fig. 4C), the pH was recovered at a similar rate compared to NBCn1 alone. Fig. 4D summarizes the mean pH$_{i}$/dt calculated during the first 2 min of recovery (n=5 for each). NBCn1 and NBCn1/PSD-95 had similar dpH$_{i}$/dt values (15.3 ± 1.7 × 10$^{-5}$ pH units/sec for NBCn1 and 14.2 ± 1.7 × 10$^{-5}$ pH units/sec for NBCn1/PSD-95; p > 0.05). These values were significantly higher than that for PSD-95 alone (3.0 ± 2.2 × 10$^{-5}$ dpH$_{i}$/dt; p < 0.05). Thus, consistent with the result from HEK 293 cells, PSD-95 has no effect on NBCn1 cotransport activity in *Xenopus* oocytes. Also, PSD-95 coimmunoprecipitated with NBCn1 in the lysates prepared from oocytes injected with NBCn1 and PSD-95, determined by coimmunoprecipitation (data not shown).

**PSD-95 increases channel-like activity of NBCn1**

We then tested whether PSD-95 affects channel-like activity of NBCn1 that produces a steady-state current in the Na/HCO$_3^-$-independent manner [2, 7]. For this analysis, I-V relationships were determined by a stair-case voltage command from –120 to 60 mV in HCO$_3^-$ /CO$_2^-$-free ND96 solution (the holding potential of –60 mV). Uninjected controls and PSD-95-injected oocytes produced negligible basal currents (Fig. 5A). In contrast, oocytes expressing NBCn1 had distinct inward currents at negative potentials and distinct outward currents at positive potentials (Fig. 5B). These are hallmarks for NBCn1 channel-like activity. The membrane potential was shifted positively (–54.1 ± 0.7 mV for controls and –25.2 ± 0.9 mV for NBCn1), another hallmark. PSD-95 increased the currents. At –54 mV, which is the average resting potential of control oocytes, the inward current of 130 nA was generated by NBCn1, while 310 nA was generated by NBCn1/PSD-95. Fig. 5C summarizes the mean slope conductance measured near the zero-current voltage. The conductance was higher when the two proteins were coexpressed (6.5 ± 0.4 µS for NBCn1 alone, n = 4 versus 11.4 ± 2.2 µS for NBCn1/PSD-95, n = 5; p < 0.05). PSD-95 coexpression caused the zero-current voltage to move positively (–25.2 ± 0.4 mV versus –18.3 ± 0.4 mV, p < 0.05), the reason of which is unclear.

**Blocking the PDZ interaction abolishes PSD-95-mediated increase in NBCn1 conductance**

To ascertain that the increased NBCn1 conductance by PSD-95 was due to their interaction, we performed two sets of experiments. In the first experiment, we deleted the last four amino acids ETSL and tested whether the deletion abolishes the PSD-95-mediated increase in NBCn1 conductance. Fig. 6A shows the results. PSD-95 increased the mean conductance for NBCn1 (3.9 ± 0.4 µS for NBCn1, n = 6 versus 6.2 ± 0.3 µS for NBCn1/PSD-95, n = 7; p < 0.05), consistent with the above finding. However, PSD-95 failed to increase the conductance when ETSL was deleted (4.0 ± 0.3 µS for NBCn1(ΔETSL)/NBCn1, n = 8). The deletion has negligible effect on NBCn1 function (data not shown), comparable with the previous report [26] that the C-terminal domain is not critical for function. In the second
experiment, we injected the peptide containing the last 15 amino acids of rat NBCn1 into oocytes expressing NBCn1 so that the peptide could compete and disrupt the interaction between the full-length NBCn1 and PSD-95. The peptide (0.05 µg/oocyte) was injected 48 h after NBCn1 and NBCn1/PSD-95 RNA injection, and then NBCn1-associated currents were measured 36 h later. Fig. 6B shows the results. In oocytes expressing NBCn1, the mean slope conductance for NBCn1 was unaffected by the peptide (2.9 ± 0.5 µS for NBCn1, n = 5 versus 2.9 ± 0.2 µS for NBCn1 with the peptide, n = 9; p > 0.05). However, in oocytes expressing NBCn1/PSD-95, the mean slope conductance was decreased by 57% by the peptide (9.8 ± 2.8 µS for NBCn1/PSD-95, n = 10 versus 4.2 ± 1.0 µS for NBCn1/PSD-95 with the peptide, n = 8; p < 0.05). Thus, the peptide competing with NBCn1 for the interaction with PSD-95 inhibited the NBCn1-mediated conductance.

Discussion

In this study, we found that PSD-95 interacts with NBCn1 and increases its channel-like activity while negligibly effecting Na/HCO₃ cotransport activity. These findings not only provide additional information on the NBCn1/PSD-95 interaction we previously reported, but they also give a new foundation for understanding the molecular and functional role of NBCn1.

Acid/base regulation is important for maintaining ionic and pH homeostasis in neurons [27], and pH disturbance is linked to neurological dysfunctions such as ischemia, coma, and epilepsy [28, 29]. Neurons possess regulatory mechanisms for maintaining pH not to fall far below the normal ranges, and among these mechanisms is Na/HCO₃ cotransport mediated by NBCn1. NBCn1 gene disruption results in severe defects in hearing and vision [5], as well as abnormal regulation of vascular tone [6], and NBCn1 knockdown reduces glutamate toxicity in neurons [30]. These reports provide strong evidence that NBCn1-mediated pH regulation play critical roles in tissues where the transporter is present [14]. Determined by electron microscopic immunoperoxidase, we demonstrate a predominant localization of NBCn1 in postsynaptic membranes (Fig. 1). Double label immunofluorescence of hippocampal neurons shows a partial colocalization of NBCn1 and PSD-95 in relatively large dendrites, consistent with the previous report [13]. The finding of NBCn1 in postsynaptic membranes brings up an interesting point on Na/HCO₃ transporters at synapses. Recent studies show that NBCn2 is present in postsynaptic membranes, whereas the Na⁺-driven Cl/HCO₃ exchanger NDCBE is located in synaptic vesicles [31]. Thus, NBCn1 and NBCn2 govern postsynaptic neuronal pH and/or synaptic cleft pH, while the Cl⁻-dependent Na/HCO₃ transporter NDCBE governs synaptic vesicular pH in the presynaptic terminal [24]. Whether NBCn2 is a Cl⁻-independent Na/HCO₃ transporter is controversial. We and our colleagues reported that NBCn2 moves
Na\(^+\) and HCO\(_3^-\), but not Cl\(^-\) [32]. The Cl\(^-\) influx caused by NBCn2 is due to Cl\(^-\) self-exchange uncoupled to Na/HCO\(_3^-\) transport. On the other hand, Damkier et al. [33] reported that NBCn2 moves Na\(^+\) and HCO\(_3^-\) in exchange for Cl\(^-\). In this case, NBCn2 is similar to NDCBE in function. Regardless, it is evident that NBCn2 has the ability to move Cl\(^-\) either directly or indirectly and is distinct from NBCn1.

The pull-down assay shows PSD-95 interaction with NBCn1, but not NBCn2 (Fig. 2A). One possible explanation is that NBCn2 variants containing the PDZ-binding motif are absent at synapse. NBCn2 immunofluorescence signals in neurons appear to be markedly diffused through the cytosol in cell bodies and dendrites [34]. Another possible explanation is that GST/NBCn2 requires a more stringent binding condition than GST/NBCn1 to pull down PSD-95 from brain lysates. Different PDZ motifs have different binding affinities of a given PDZ domain, and an optimal binding depends upon in vitro conditions such as ion stringency and temperature [24]. The difference between NBCn1 and NBCn2 is the second last amino acid in the PDZ motif (ETSL vs. ETCL). The second last residue is known to be least critical for PDZ-mediated interaction, but some PDZ domains including PSD-95 can distinguish amino acids at this position [24].

The interaction between NBCn1 and PSD-95 is confirmed by coimmunoprecipitation in HEK 293 cells (Fig. 2B). This interaction occurs without influencing PSD-95’s essential ability to interact with NMDA receptors via the first and second PDZ domains [14]. The binding domain for NBCn1 is less clear although our preliminary proteomic analysis shows the third PDZ domain of PSD-95 being responsible for NBCn1 interaction (unpublished observation). Thus, NBCn1 and NR2A do not share the binding domains in PSD-95. In this sense, there is a possibility that NBCn1 clusters with NMDA receptors via PSD-95 to form a ternary complex containing NBCn1/PSD-95/NR2A although it is unclear if this complex exists in neurons.
Nonetheless, we do not exclude another possibility that our coimmunoprecipitation data could be due to partial expression of the three proteins in transfected cells. NBCn1/PSD-95 interaction would not affect NR2A/PSD-95 interaction if some cells express PSD-95 and NBCn1 while others express PSD-95 and NR2A. Regarding the binding strength, we observe that a large fraction of NR2A proteins in the lysates bind to PSD-95, while a small fraction of NBCn1 proteins bind to PSD-95. It could be due to either different binding affinities between the two interactions or different numbers of PSD-95 PDZ domains for NBCn1 or NR2A. A stronger intensity does not necessarily reflect a biologically more significant role of the corresponding interaction, although we believe that PSD-95/NR2A interaction would be more functionally significant than NBCn1/PSD-95 interaction.

Our pH experiments reveal no significant difference in dpH/dt between NBCn1 and NBCn1/PSD-95 in both HEK 293 cells and Xenopus oocytes. Thus, PSD-95 has negligible effect on NBCn1 cotransport regardless of two different expression systems. On the other hand, PSD-95 increases the transporter’s channel-like activity that produces a steady-state inward current at the resting voltage (Fig. 5). PSD-95 increases the slope conductance without significantly altering the zero-current voltage, implying that more pores are opened by PSD-95. Deleting the C-terminal ETSL abolishes the stimulatory effect of PSD-95, and the peptide containing ETSL competes with the full-length transporter (Fig. 6). Together, these results strongly support the idea that PSD-95 binds to the transporter and stimulates its channel-like activity. We propose that the channel-like activity may occur via an intermolecular cavity of multimeric NBCn1 proteins. NBCe1 (SLC4A4), which has high amino acid sequence homology with NBCn1, forms a dimer consisting of two individually functional subunits [35, 36]. The distantly related bicarbonate transporters AE1 (SLC4A1) and AE2 (SLC4A2) are also homodimers [37]. We believe that NBCn1 is also in a dimeric form with each functional subunit, and PSD-95 may facilitate the dimerization process. We also note that PSD-95 is a dimer [38]. A similar example of PDZ protein-mediated intermolecular modification is found in CFTR [43]. The PDZ protein CAP70 (PDZK1) enhances intermolecular CFTR-CFTR contact and switches CFTR channels to a more active conducting state.

Regardless of the underlying mechanism, PSD-95 substantially enhances NBCn1-mediated steady state currents (Fig. 5) that raise intracellular Na⁺ levels in mammalian cells and frog oocytes [2, 7]. The increased Na⁺ levels would lower an electrochemical Na⁺ gradient across cell membranes and reduce a driving force of sodium channels/transporters if they are present with NBCn1 in the same cells. This inhibitory effect would be augmented by PSD-95. It is important to note that the channel-like activity of NBCn1 has been characterized so far in heterologous overexpression systems, which are different from the native system. It is unclear whether NBCn1 has a similar activity and is stimulated by PSD-95 in neurons, where the interaction of NBCn1 and PSD-95 depends upon the spatial distribution and expression level of the two proteins. PSD-95 binds to a large number of different postsynaptic proteins and forms a constituent of macromolecular complexes. Also, NBCn1 is expected to bind to different PDZ proteins via its C-terminal PDZ motif. It is likely that the interaction is very limited and local in neurons.

NBCn1 is not the only HCO₃⁻ transporters exhibiting channel-like activity, and other transporters have similar properties. AE1 in erythrocytes mediates a conductive anion flux that is DIDS-insensitive and increases with increasing membrane hyperpolarization [39]. The Cl/HCO₃ exchangers Slc26a3 (DRA, CLD) and Slc26a6 (CFEX, PAT-1) mediate large NO₃⁻ and SCN⁻ currents uncoupled to OH⁻ or HCO₃⁻ transport [40, 41]. The electroneutral NDAE1 in Drosophila has a small inward current [42]. The functional implication of these conductance properties is unclear.

In summary, our study shows that PSD-95 binds to NBCn1 via the C-terminal PDZ motif and alters NBCn1-mediated conductance, but not Na/HCO₃ cotransport. To our knowledge, these findings indicate for the first time that PSD-95 separates NBCn1’s dual activities. It will be interesting to investigate whether a similar mechanism is found in neurons in future experiments.
Acknowledgements

We thank Jeff Pare and Susan Jenkins for technical assistance with electron microscopy procedures, Drs. Suzanne Zukin (Albert Einstein College) and Makato Inui (Yamaguchi University) for providing PSD-95, and Drs. Randy Hall and Steve Traynelis (Pharmacology, Emory) for providing PSD-95, NR1A, and NR2A plasmids. This work was supported by grants from the NIH GM078502, American Heart Association, and Emory URC (I. C.), Yerkes National Primate Center NIH base grant RR00165 (Y. S.), and DK061418 (C.C.Y.).

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