Presynaptic Release Probability Is Increased in Hippocampal Neurons From ASIC1 Knockout Mice

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Cho J-H, Askwith CC. Presynaptic release probability is increased in hippocampal neurons from ASIC1 knockout mice. J Neurophysiol 99: 426–441, 2008. First published December 19, 2007; doi:10.1152/jn.00940.2007. Acid-sensing ion channels (ASICs) are voltage-independent cation-permeable ion channels activated by extracellular acidosis (Krishtal 2003; Waldmann 2001; Wemmie et al. 2006). These H+-gated channels are members of the degenerin/epithelial Na+ (DEG/ENaC) family (Kellenberger and Schild 2002). Four ASIC genes encode at least six subunits (ASIC1a, 1b, 2a, 2b, 3, and 4) through alternative splicing. ASIC subunits are expressed in central and peripheral neurons, and the characteristics of H+-gated currents are determined by the ASIC subunits expressed within the cell. ASIC1a, ASIC2a, and ASIC2b are expressed throughout the brain with particularly high abundance in the cerebral cortex, hippocampus, amygdala, olfactory bulb, and the cerebellum (García-Anoveros et al. 1997; Linguegila et al. 1997; Price et al. 1996; Waldmann et al. 1996, 1997). ASIC1a forms not only heteromeric channels with ASIC2a, but also Ca2+-permeable homomeric channels (Askwith et al. 2004; Bassilana et al. 1997; Benson et al. 2002; Gao et al. 2007; Hesselager et al. 2004; Yermolaieva et al. 2004). ASIC1a is enriched in synaptosomal fractions and expressed in dendritic spines (Hruska-Hageman et al. 2002; Wemmie et al. 2002). ASIC1 knockout mice display impaired spatial learning, eye-blink conditioning, and fear conditioning (Wemmie et al. 2002, 2003). In turn, transgenic mice overexpressing ASIC1a exhibit enhanced fear conditioning (Wemmie et al. 2004). Together, these observations suggest that ASIC1a plays a role in synaptic transmission.

Wemmie et al. (2002) performed extracellular field potential recordings of hippocampal slices and observed that hippocampal CA1 long-term potentiation (LTP) was impaired in ASIC1 knockout mice. Specifically, N-methyl-D-aspartate receptor (NMDAR) activation during high-frequency stimulation was reduced in hippocampal CA3–CA1 synapses of ASIC1 knockout mice. The authors hypothesized that protons released from synaptic vesicles activate postsynaptic ASICs, which depolarize the postsynaptic membrane and facilitate activation of NMDARs by relieving Mg2+ block (Wemmie et al. 2002). Depolarization-induced removal of Mg2+ block from the NMDAR is usually mediated by the activation of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptors (AMPARs). In specific situations, such as silent synapses that express few functional AMPARs in the postsynaptic membrane, ASICs might induce NMDAR activation and thereby facilitate LTP. ASIC-mediated H+-gated current, however, has not yet been observed during high-frequency stimulation, and inhibition of ASIC channels does not appear to affect postsynaptic currents (Alvarez de la Rosa et al. 2003). These results suggest that ASICs may influence synaptic transmission through an alternate mechanism.

In this study, we examined the role of ASICs in synaptic transmission using microisland cultures of hippocampal neurons from wild-type and ASIC knockout mice. Solitary neurons in microisland culture are synaptically isolated from other neurons; monosynaptic connections (autapses) form between the axon and dendrites of the same neuron (Bekkers and Stevens 1991). This preparation allows a detailed analysis of presynaptic and postsynaptic parameters of synaptic transmission in the absence of complex polysynaptic circuitry (Bekkers and Stevens 1991). Microisland cultures have been widely used for electrophysiologic studies of synaptic transmission on both cellular and molecular levels (Chavis and Westbrook 2001; Rhee et al. 2002; Rosenmund and Stevens 1996; Wierda et al. 2007). Using this method, we determined that the prob-
ability of neurotransmitter release is increased in neurons from ASIC1 knockout mice, suggesting that ASIC1a can influence glutamatergic synaptic transmission through presynaptic mechanisms.

METHODS

Microisland culture of hippocampal neurons

Primary hippocampal neuron cultures were prepared using previously published methods (Askwith et al. 2004; Cho and Askwith 2007). Briefly, hippocampi were dissected from postnatal day (P) 0–P1 pups, freed from extraneous tissue, and cut into pieces. ASIC1 and ASIC2 knockout mice develop and breed normally, and there are no overt abnormalities in brain morphology (Price et al. 2000; Wemmie et al. 2002). Hippocampal tissue was transferred into Leibovitz’s L-15 medium containing 0.25 mg/ml bovine serum albumin and 0.38 mg/ml papain, and incubated for 15 min at 37°C with 95% O2-5% CO2 gently blown over the surface of the medium. After incubation, the hippocampal tissue was washed three times with mouse M5-5 medium (Earle’s minimal essential medium with 5% fetal bovine serum, 5% horse serum, 0.4 mM L-glutamine, 16.7 mM glucose, 5,000 U/l penicillin, 50 mg/l streptomycin, 16 mM selenite, 1.4 mg/l transferrin) and triturated. For conventional mass cultures, a collagen solution containing 0.5 mg/ml rat tail collagen in 1:1,000 acetic acid was spread onto 10-mm glass coverslips. For microisland cultures, a microatomizer was used to spray a fine mist of collagen solution onto the coverslips (Bekkers and Stevens 1991). The collagen was allowed to dry completely and then the coverslips were exposed to UV light for 1 h. Hippocampal cells were plated on coverslips in 24-well dishes at a density of 500,000 cells per well for mass culture and 250,000 cells per well for microisland culture. After 48–72 h, 10 μM cytosine β-d-arabinofuranoside was added to inhibit glial proliferation. Neurons in conventional mass culture were used from 9 to 13 days in culture and neurons in microisland culture were used from 12 to 19 days in culture. Multiple breeding pairs of ASIC1 knockout (three pairs), ASIC2 knockout (three pairs), and wild-type (composed of three pairs siblings of ASIC2 knockout) mice were used to produce neonatal pups. No difference was observed in H+–gated currents or postsynaptic responses between wild-type neurons from the ASIC1 or ASIC2 knockout mouse lines.

Transfection of primary hippocampal neurons

Human ASIC1a cDNA (GenBank Accession Number NM_001095) was cloned into the pcDNA3.1 expression vector (Invitrogen, Carlsbad, CA). Hippocampi were dissected from ASIC1 knockout pups of P0–P1 as before except that hippocampal cells were transfected with the ASIC1a or vector just prior to plating. Briefly, dissociated hippocampal cells were split into two groups from the same preparation (3–4 million cells each group), suspended in 100 μl of Nucleofector Solution from the Basic Nucleofector Kit (Amaxa, Gaithersburg, MD), and mixed with 1 μg of pEGFP-C1 (Clontech, Mountain View, CA) as well as 2 μg of either the ASIC1α construct or vector alone (pcDNA3.1). Hippocampal cells were electroporated with the Nucleofector II (program O-05, Amaxa) and plated on collagen-spritzed coverslips in 24-well dishes at a density of 500,000–1,000,000 cells per well in M5-5 media. Culture medium was replaced with fresh M5-5 media at a density of 500,000 cells per well for microisland cultures, a microatomizer was used to spray a fine mist of M5-5 media (Earle’s minimal essential medium with 5% fetal bovine serum, 5% horse serum, 0.4 mM L-glutamine, 16.7 mM glucose, 5,000 U/l penicillin, 50 mg/l streptomycin, 16 mM selenite, 1.4 mg/l transferrin) and triturated. For conventional mass cultures, a collagen solution containing 0.5 mg/ml rat tail collagen in 1:1,000 acetic acid was spread onto 10-mm glass coverslips. For microisland cultures, a microatomizer was used to spray a fine mist of collagen solution onto the coverslips (Bekkers and Stevens 1991). The collagen was allowed to dry completely and then the coverslips were exposed to UV light for 1 h. Hippocampal cells were plated on coverslips in 24-well dishes at a density of 500,000 cells per well for mass culture and 250,000 cells per well for microisland culture. After 48–72 h, 10 μM cytosine β-d-arabinofuranoside was added to inhibit glial proliferation. Neurons in conventional mass culture were used from 9 to 13 days in culture and neurons in microisland culture were used from 12 to 19 days in culture. Multiple breeding pairs of ASIC1 knockout (three pairs), ASIC2 knockout (three pairs), and wild-type (composed of three pairs siblings of ASIC2 knockout) mice were used to produce neonatal pups. No difference was observed in H+–gated currents or postsynaptic responses between wild-type neurons from the ASIC1 or ASIC2 knockout mouse lines.

Electrophysiology

To record H+–gated and postsynaptic currents, we used the whole cell voltage-clamp technique. The extracellular solution contained 128 mM NaCl, 5.4 mM KCl, 2 mM CaCl2, 1 mM MgCl2, 5.55 mM glucose, 1 μM glycine, and 0.8 mM HEPES. Unless otherwise indicated, this low concentration of pH buffer (HEPES) was used to facilitate ASIC activation by endogenous acidic fluctuations in pH. Tetramethylammonium hydroxide was used to adjust the pH of the extracellular solution to either pH 7.4 or pH 6.0. An extracellular pH 6.0 solution containing 10 mM HEPES and 10 mM MES as pH buffers was used to evoke H+–gated currents. The intracellular pipette solution contained 121 mM KCl, 10 mM NaCl, 2 mM MgCl2, 5 mM EGTA, 10 mM HEPES, 2 mM Mg-ATP, and 300 μM Na2-GTP (pH 7.25). For perforated-patch recording, we used the intracellular pipette solution containing 130 mM K-glucionate, 20 mM KCl, 10 mM HEPES, and 0.1 mM EGTA (pH 7.3). The pipette tip was filled with the intracellular solution and back-filled with the solution containing 150 μg/ml of nystatin. Patch electrodes were pulled with a P-97 micropipette puller (Sutter Instrument, Novato, CA) and fire-polished with a microforge (Narishige, East Meadow, NY). Micropipettes with 2–4 MΩ were used for experiments. The membrane potential was held constant at −70 mV. Data were collected at 5 kHz using an Axopatch 200B amplifier, Digidata 1322A, and Clampex 9 (Molecular Devices, Sunnyvale, CA). Neurons were continuously superfused with the extracellular solution from gravity-fed perfusion pipes at a flow rate of about 1 ml/min. Perfusion pipes were placed 250 to 300 μm away from cells, and flow was directed toward the recorded cells to ensure fast solution exchange. There was no significant difference in whole cell membrane capacitance of neurons from the three genotypes (95 ± 10, 101 ± 17, 85 ± 10 pF, n = 24, 10, 22 for wild-type, ASIC1 knockout, and ASIC2 knockout neurons, respectively), suggesting neuron size was not different between genotypes.

H+–gated currents were evoked by the exogenous application of pH 6.0 extracellular solutions. Desensitization time constant (τs) was calculated by fitting the decay phase of H+–gated currents to the single-exponential equation, \( I(t) = I_{\text{max}} \exp(-t/\tau_s) \), where \( I_{\text{max}} \) is the peak amplitude of H+–gated current at pH 6.0. To record action potential (AP)–evoked whole cell postsynaptic currents, solitary autaptic neurons on microislands were selected and stimulated every 10 s (0.1 Hz) with a transient depolarization of membrane potential from −70 to +10 mV for 2 ms. Extracellular solution containing 10 mM 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX) / 50 μM d-2-amino-5-phosphonovaleric acid (AP5) or 30 μM bicuculline was used to determine whether the autaptic neuron was glutamatergic or GABAergic. To isolate AMPAR- or NMDAR-mediated excitatory postsynaptic current (EPSC), we used 1 mM Mg2+ or 10 μM CNQX in Mg2+-free solution, respectively. We attained similar results when the AMPAR EPSC was isolated with 50 μM AP5. Decay time constants (τ) of AMPAR EPSC or GABA\textsubscript{A} receptor-mediated postsynaptic current (GABA\textsubscript{A}R PSC) were calculated by fitting curves of postsynaptic currents to the single-exponential equation, \( I(t) = I_{\text{max}} \exp(-t/\tau) \), where \( I_{\text{max}} \) is the peak amplitude of AMPAR EPSC or GABA\textsubscript{A}R PSC. To quantify the decay rate of NMDAR...
EPSCs, we calculated the weighted mean decay time constant of NMDAR EPSC. NMDAR EPSC curves were fitted to the double-exponential equations, \( I(t) = I_1 \exp(-t/\tau_1) + I_2 \exp(-t/\tau_2) \), where \( \tau_1 \) and \( \tau_2 \) are fast and slow components of peak amplitudes and decay time constants of NMDAR EPSC; the weighted mean decay time constants (\( \tau_{\text{w}} \)) were calculated from the equation, 
\[
\tau_{\text{w}} = \frac{\tau_1 I_1 + \tau_2 I_2}{I_1 + I_2}.
\]
Miniature EPSC (mEPSC) and hypertonic (500 mM) sucrose-induced currents were recorded from neurons in conventional mass culture using the extracellular solution containing 1 mM Mg\(^{2+}\), 30 \( \mu \)M bicuculline, and 1 \( \mu \)M tetrodotoxin. Unless otherwise indicated, all reagents were purchased from Invitrogen/Gibco (Carlsbad, CA), Sigma–Aldrich (St. Louis, MO), or Fisher Scientific (Waltham, MA).

Data were analyzed using Clampfit 9 software (Molecular Devices). We analyzed mEPSC using the template search function of Clampfit 9. Data are presented as means ± SE. As appropriate, a two-tailed Student’s \( t \)-test and one-way ANOVA with Bonferroni’s simultaneous multiple comparisons were used for statistical analyses and performed with Minitab15 software (Minitab, State College, PA).

**RESULTS**

**\( H^+ \)-gated currents are altered in ASIC knockout neurons cultured on microislands**

In conventional mass-cultured hippocampal neurons, ASIC1a homomultimers and ASIC1a/ASIC2a heteromultimers are responsible for the majority of \( H^+ \)-gated currents (Askwith et al. 2004). The ASIC1a subunit plays a dominant role, and loss of ASIC1a eliminates \( H^+ \)-gated currents evoked by pH values >5 (Wemmie et al. 2002; Xiong et al. 2004). The ASIC2a subunit modulates \( H^+ \)-gated current by forming heteromultimeric channels with ASIC1a that desensitize more rapidly and recover from previous acid applications more quickly compared with homomultimeric ASIC1a channels (Askwith et al. 2004; Benson et al. 2002). To determine whether ASIC1 and ASIC2 have similar roles in microisland culture, we analyzed \( H^+ \)-gated currents of solitary hippocampal neurons from wild-type, ASIC1 knockout, and ASIC2 knockout mice. Application of acidic extracellular solution (pH 6.0) did not induce a substantial transient current in ASIC1 knockout neurons (Fig. 1, A and B, peak amplitude = 39 ± 17 pA, \( n = 13 \), \( P < 0.01 \), wild-type vs. ASIC1 knockout, one-way ANOVA). By comparison, extracellular application of pH 6.0 solution induced transient inward currents in both wild-type and ASIC2 knockout neurons (Fig. 1, A and B, peak amplitude = 796 ± 106 pA, \( n = 53 \) for wild-type, and 1,595 ± 217 pA, \( n = 31 \) for ASIC2 knockout). The peak amplitude and desensitization time constant (\( \tau_d \)) of \( H^+ \)-gated currents were significantly larger in ASIC2 knockout neurons compared with wild-type neurons (Fig. 1, B and C, \( P < 0.01 \) for peak amplitude, one-way ANOVA; \( P < 0.05 \) for desensitization time constants, unpaired \( t \)-test). Recovery from desensitization was also substantially slower in ASIC2 knockout neurons compared with wild-type neurons (Fig. 1, A and D, \( 45 ± 3% \), \( n = 37 \) for wild-type, 17 ± 3%, \( n = 31 \) for ASIC2 knockout, \( P < 0.0001 \), unpaired \( t \)-test). These results are consistent with the loss of ASIC1a/ASIC2a heteromeric channels, which desensitize and recover faster, in the ASIC2 knockout neurons (Askwith et al. 2004; Benson et al. 2002). These results

**FIG. 1.** \( H^+ \)-gated currents of autaptic neurons in microisland cultures. A: representative traces of \( H^+ \)-gated currents of wild-type, acid-sensing ion channel 1 knockout (ASIC1 KO) and ASIC2 knockout (ASIC2 KO) neurons in microisland culture. To evoke \( H^+ \)-gated currents, we exchanged the extracellular solution of pH 7.4 to pH 6.0 for 6–7 s (gray bars). To estimate recovery from desensitization, the pH of the extracellular solution was returned to 7.4 for about 2.5 s, and then a second pH 6.0 application was made in wild-type and ASIC2 knockout neurons. B: peak amplitude of pH 6.0–induced \( H^+ \)-gated currents. C: the desensitization time constants (\( \tau_d \)) of \( H^+ \)-gated current at pH 6.0. D: recovery from desensitization was determined by applying 2 pulses of pH 6.0 with an interval of 2.5 s. Recovery was assessed by comparing the peak current amplitude in response to the first pH 6.0 applications to the peak current amplitude of the second pH 6.0 application. E: the current density of \( H^+ \)-gated currents in glutamatergic and GABAergic autaptic neurons from wild-type mice. Current density was calculated by dividing the peak amplitude of pH 6.0–induced current by whole cell membrane capacitance. Numbers in parentheses indicate the numbers of neurons examined. All data in this and subsequent figures are expressed as means ± SE. One-way ANOVA (B) and unpaired \( t \)-test (C–E) were used to assess statistical significance.
indicate that ASIC subunits play similar roles in microisland culture and conventional mass culture (Askwith et al. 2004). Whether a solitary neuron in microisland culture is glutamatergic or GABAergic can be determined by analyzing the AP-evoked postsynaptic current. Extracellular solutions containing CNQX/AP5 or bicuculline were used to identify the autaptic neuron as glutamatergic or GABAergic, respectively. Both glutamatergic and GABAergic autaptic neurons expressed H^+ -gated currents in wild-type and ASIC2 knockout neurons. The current density (peak current amplitude divided by whole cell membrane capacitance) of H^+ -gated currents was significantly greater in GABAergic neurons than in glutamatergic neurons (Fig. 1B, 13.7 ± 2.2 pA/pF, n = 24 for glutamatergic neurons, and 22.3 ± 2.7 pA/pF, n = 18 for GABAergic neurons, P < 0.05, unpaired t-test). The desensitization time constant and recovery from desensitization of H^+ -gated currents were not different between these two populations of neurons (data not shown). Together, these data indicate that both glutamatergic and GABAergic neurons in microisland culture express H^+ -gated channels with biophysical characteristics similar to those of mass-cultured neurons.

Whole cell postsynaptic currents in solitary neurons in microisland culture

To gain insight into the role of ASICs in synaptic transmission, AP-evoked postsynaptic currents of solitary neurons in microisland culture were analyzed using the whole cell voltage-clamp technique. In GABAergic autaptic neurons there was no significant difference in the peak amplitude or decay time constant of GABAAR-mediated postsynaptic current between wild-type and ASIC knockout neurons (Fig. 2, A and B, peak amplitude = 4.19 ± 0.43, 3.48 ± 0.34, 4.52 ± 0.56 nA, n = 61, 48, 32 for wild-type, ASIC1 knockout, and ASIC2 knockout, respectively, P = 0.27; decay time constant = 119 ± 24, 132 ± 27, 115 ± 22 ms, n = 12, 14, 13 for

![Graph showing AMPAR and NMDAR EPSCs](image)

FIG. 2. Whole cell action potential (AP)–evoked postsynaptic currents in GABAergic and glutamatergic autaptic neurons. A: representative traces of whole cell GABA_A receptor-mediated postsynaptic current (GABA_A PSC) of wild-type, ASIC1 knockout, and ASIC2 knockout neurons. 30 ms depolarization to +10 mV from a −70-mV holding potential was used to evoke neurotransmitter release and postsynaptic currents. B: quantification of peak amplitude of GABA_A PSC. n = 62, 48, and 32 for wild-type, ASIC1 knockout, and ASIC2 knockout. C: representative traces of whole cell AMPAR–mediated excitatory postsynaptic current (EPSC, black), and NMDAR–mediated EPSC (gray) of wild-type, ASIC1 knockout, and ASIC2 knockout neurons. AMPAR EPSCs were recorded in the presence of 1 mM Mg^2+. NMDAR EPSCs were isolated in Mg^2+-free extracellular solution using 10 μM 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX), an AMPAR antagonist. Membrane potential was held at −70 mV. D–F: average peak amplitude of the AMPAR EPSCs (D), NMDAR EPSCs (E), and the average AMPAR EPSC/NMDAR EPSC ratio from individual neurons (F). n = 40, 42, and 28 for wild-type, ASIC1 knockout, and ASIC2 knockout. One-way ANOVA was used to assess statistical significance.
wild-type, ASIC1 knockout, and ASIC2 knockout, respectively, \( P = 0.87 \), one-way ANOVA). In glutamatergic autaptic neurons, AMPAR- and NMDAR-mediated EPSCs were individually isolated using either 1 mM Mg\(^{2+}\) or 10 \( \mu \)M CNQX, respectively (Fig. 2C). Although there was a slight increase in the average peak amplitudes in ASIC1 knockout neurons, there was no significant difference between wild-type and ASIC knockout neurons in the peak amplitudes of either the AMPAR EPSC or NMDAR EPSC (Fig. 2, D and E, peak amplitude of AMPAR EPSC = 3.65 \( \pm \) 0.50, 4.29 \( \pm \) 0.58, 3.87 \( \pm \) 0.60 nA, \( n = 40, 42, 28 \) for wild-type, ASIC1 knockout, and ASIC2 knockout, respectively, \( P = 0.69 \); peak amplitude of NMDAR EPSC = 1.10 \( \pm \) 0.17, 1.58 \( \pm \) 0.26, 1.08 \( \pm \) 0.18 nA for wild-type, ASIC1 knockout, and ASIC2 knockout respectively, \( P = 0.17 \), one-way ANOVA). There was also no significant difference in the decay time constant of AMPAR or NMDAR EPSC (decay time constant of AMPAR EPSC = 11.4 \( \pm \) 0.8, 13.3 \( \pm \) 1.3, 10.6 \( \pm \) 0.9 ms, \( n = 36, 40, 27 \) for wild-type, ASIC1 knockout, and ASIC2 knockout respectively, \( P = 0.18 \); weighted mean decay time constant of NMDAR EPSC = 198 \( \pm \) 14, 182 \( \pm \) 9, 155 \( \pm \) 11 ms, \( n = 45, 48, 28 \) for wild-type, ASIC1 knockout, and ASIC2 knockout, respectively, \( P = 0.07 \), one-way ANOVA). To compare the relative contribution of AMPARs and NMDARs to the total EPSC, we recorded the AMPAR/NMDAR EPSC ratio was significantly smaller in both AMPAR and NMDAR EPSCs in the same neurons and of AMPARs and NMDARs to the total EPSC, we recorded in ASIC1 knockout neurons (Fig. 2, B). H\(^+\)-dependent current and thus display a smaller AMPAR/NMDAR EPSC ratio. To test this hypothesis, we examined whether ASICs directly contribute to postsynaptic currents. First, we determined whether ASIC current could be isolated during synaptic transmission. Both AMPAR and NMDAR EPSCs were simultaneously inhibited using 10 \( \mu \)M CNQX and 50 \( \mu \)M AP5 in wild-type glutamatergic neurons on microislands. When both AMPARs and NMDARs were blocked, a small transient residual current was observed in glutamatergic neurons that contributed to the peak AMPAR EPSC (Fig. 3A). This current was not significantly different between wild-type and ASIC1 knockout neurons (128 \( \pm \) 31 pA, \( n = 7 \) for wild-type neurons; 124 \( \pm \) 50 pA, \( n = 7 \) for ASIC1 knockout neurons, \( P = 0.96 \), unpaired \( t \)-test). We also recorded both the AMPAR and the residual current in the same neuron and calculated the ratio of the residual current amplitude to the peak amplitude of the AMPAR EPSC. The ratio was not different between wild-type and ASIC1 knockout neurons (Fig. 3B, 4.06 \( \pm \) 0.77\%, \( n = 7 \) for wild-type neurons; 5.18 \( \pm \) 0.96\%, \( n = 7 \) for ASIC1 knockout neurons, \( P = 0.38 \), unpaired \( t \)-test), suggesting that the contribution of the residual current to peak AMPAR EPSC was not affected by the loss of ASIC1.

We also examined the effect of amiloride (300 \( \mu \)M), a nonspecific ASIC blocker, on residual current and AMPAR EPSCs (Fig. 3, C and E). H\(^+\)-gated currents evoked by exogenous acid application were blocked nearly completely by 300 \( \mu \)M amiloride (Fig. 3C). However, the contribution of the residual current to AMPAR EPSCs in wild-type glutamatergic neurons was not affected by amiloride (Fig. 3, C and D, 3.62 \( \pm \) 0.80 and 3.30 \( \pm \) 0.98\% before and during amiloride application, respectively, \( n = 5 \), \( P = 0.38 \), paired \( t \)-test). Furthermore, AMPAR EPSCs were not altered by amiloride in either wild-type or ASIC1 knockout neurons (Fig. 3, E and F, 92.1 \( \pm \) 4.1\% of pretreatment control, \( n = 8 \), \( P = 0.10 \), paired \( t \)-test for wild-type; 99.7 \( \pm \) 4.6\%, \( n = 8 \), \( P = 0.35 \), paired \( t \)-test for ASIC1 knockout; \( P = 0.23 \), wild-type vs. ASIC1 knockout, unpaired \( t \)-test). Together, these results indicate that ASIC currents are not components of the EPSC and the altered AMPAR/NMDAR EPSC ratio in ASIC1 knockout neurons cannot be explained by loss of postsynaptic ASIC currents evoked during synaptic transmission. Thus another mechanism is responsible for the decrease in AMPAR/NMDAR EPSC ratio in the ASIC1 knockout neurons.

### Exogenous acid application did not alter whole cell postsynaptic currents

Activation of ASIC1a channels could induce changes in AMPAR-, NMDAR-, or GABA\(_A\)R-mediated postsynaptic cur-

**ASIC currents are not components of EPSCs**

If ASIC currents contribute directly to the postsynaptic current, then loss of H\(^+\)-gated currents could explain the altered AMPAR/NMDAR EPSC ratio in the ASIC1 knockout neurons. ASIC1a is enriched in synaptosomal fractions and expressed in dendritic spines, suggesting ASICs play a role at postsynaptic sites (Hruska-Hageman et al. 2002; Wemmie et al. 2002, 2004). The pH inside synaptic vesicles is acidic (pH < 6.0) (Miesenbock et al. 1998) and protons are released from synaptic vesicles during synaptic transmission. Under certain circumstances, such as high-frequency synaptic transmission, these released protons can lower the pH of the synaptic cleft (Krishal et al. 1987) and affect pH-sensitive ion channel activity. This has been shown for voltage-gated Ca\(^{2+}\) channels in retinal cone photoreceptor and bipolar cells (DeVries 2001; Hosoi et al. 2005; Palmer et al. 2003; Vessey et al. 2005). ASIC activation evoked by protons released from synaptic vesicles has not been demonstrated, although spontaneous autocrine release of protons activates endogenous ASIC currents in HEK293 cells (Lalo et al. 2007). Protons released from synaptic vesicles may be rapidly trapped by endogenous pH buffers, and the acute activation of ASICs by protons would likely be for only a short time (Kristal et al. 1987). Thus protons released during synaptic transmission may activate ASIC1a-containing channels, generating postsynaptic currents in wild-type neurons, which could contribute to the early component of EPSC (AMPAR EPSC). ASIC1 knockout neurons would lack this H\(^+\)-dependent current and thus display a smaller AMPAR/NMDAR EPSC ratio. To test this hypothesis, we examined whether ASICs directly contribute to postsynaptic currents.
rents through signal transduction cascades. To test this, we measured whole cell postsynaptic currents before and after activating ASICs with exogenously applied acid. Postsynaptic currents were recorded every 10 s in wild-type autaptic neurons. After the peak amplitude of baseline postsynaptic currents stabilized, we briefly evoked H⁺-gated currents with the application of pH 6.0 extracellular solutions for 5 s. We then returned the pH to 7.4 and continued to measure postsynaptic currents every 10 s for several minutes (Fig. 4). Peak amplitudes of AMPAR, NMDAR, and GABA_A receptors were unchanged following acid application (Fig. 4, A–C, n = 4–5). AMPAR EPSCs were increased robustly when we applied 2 μM 4-β-phorbol-12,13-dibutyrate (PDBu), a diacylglycerol analog that potentiates synaptic transmission through protein kinase C and other mechanisms (Wierda et al. 2007) (Fig. 4D, n = 6). This indicates that our experimental configuration could support signal transduction–dependent changes in synaptic transmission. To further conserve intracellular signaling mechanisms, we performed the same experiment using nystatin-based perforated-patch recording. Again, we did

**FIG. 3.** ASIC currents do not contribute to the peak amplitude of EPSC in glutamatergic autaptic neurons. A: representative traces of AMPAR and NMDAR EPSC (gray traces) of a wild-type autaptic neuron. In the presence of 10 μM CNQX and 50 μM t-2-aminophosphonovaleric acid (AP5), small residual current was observed (black trace). Vertical dotted lines indicate the time of the peak amplitude of AMPAR and NMDAR EPSCs. A voltage step above the traces indicates transient membrane depolarization from −70 to 10 mV for 2 ms used to evoke an AP. B: the ratio of the residual current measured in the presence of CNQX and AP5 to the peak AMPAR EPSC in wild-type (n = 7) and ASIC1 knockout (n = 7) neurons. There is no significant difference in the ratio of residual current to AMPAR EPSC between wild-type and ASIC1 knockout neurons (unpaired t-test). C: representative traces of AMPAR EPSCs and residual current in the presence of CNQX and AP5 in a wild-type autaptic neuron (top traces). AMPAR EPSCs (a, gray trace) were recorded in the presence of AP5. Residual current (b, black trace) was isolated using CNQX and AP5 and was not affected by 300 μM amiloride (c, gray trace). Bottom traces show amiloride (300 μM) inhibition of H⁺-gated currents evoked by extracellular acid application (pH 6.0, gray bars). D: the ratio of the residual current to the AMPAR EPSC before and during amiloride application. Amiloride did not change the ratio significantly (n = 5, paired t-test). E: representative traces of AMPAR EPSCs before (−amiloride, black traces) and during 300 μM amiloride application (+amiloride, gray traces) in wild-type and ASIC1 knockout neurons. F: quantification of the effect of amiloride on AMPAR EPSCs. Current amplitude during amiloride application (+amiloride) was normalized to current amplitude before amiloride treatment (−amiloride). Amiloride did not change the peak amplitude of AMPAR EPSC significantly (n = 8, P = 0.10 for wild-type, and n = 8, P = 0.65 for ASIC1 knockout, paired t-test).
Paired-pulse ratio is reduced in ASIC1 knockout neurons

Our results suggest that the decrease in the AMPAR/ NMDAR EPSC ratio in ASIC1 knockout neurons is not due to activation of postsynaptic ASIC currents during the recording interval. To determine whether other aspects of synaptic transmission are altered in ASIC knockout neurons, we assessed the synaptic response to repetitive and paired-pulse stimulation (Mennerick and Zorumski 1995). A pair of stimuli with short intervals from 20 to 400 ms was applied, and the paired-pulse ratio (PPR) was determined by dividing the peak amplitude of the second AMPAR EPSC by the peak amplitude of the first EPSC (Fig. 5A). Compared with wild-type neurons, the PPRs with interpulse intervals of 50, 100, and 200 ms were significantly reduced in ASIC1 knockout neurons (Fig. 5, A and B). The PPR was reduced. The reason for this discrepancy is unclear. However, more variables such as polysynaptic activity contribute to the PPR in hippocampal slices compared with the PPR in microisland cultures (Mennerick and Zorumski 1995), and the simplicity and enhanced experimental control over these variables in the microisland culture system may have facilitated the detection of alterations in PPR. However, we also tested whether our pH buffer conditions may have allowed observation of altered PPR. Our experiments use a lower HEPES concentration (0.8 mM) compared with others (10 mM) to facilitate ASIC activation by endogenous acidic fluctuations in pH. To determine whether the difference in PPR was due to the low concentration of pH buffer used in our studies, we analyzed postsynaptic currents and PPR when the external buffer concentration (0.8 mM) compared with others (10 mM) to facilitate ASIC activation by endogenous acidic fluctuations in pH. To determine whether the difference in PPR was due to the low concentration of pH buffer used in our studies, we analyzed postsynaptic currents and PPR when the external buffer concentration was increased to 10 mM HEPES, a concentration of HEPES under 5% CO2). We found that increasing the concentration of HEPES from 0.8 to 10 mM slightly reduced the peak amplitude of AMPAR EPSCs, NMDAR EPSCs, and GABAAR PSCs. However, this effect was not different between wild-type, ASIC1 knockout, and ASIC2 knockout neurons (Fig. 6, A and B).
AMPAR EPSC, NMDAR EPSC, and GABA<sub>A</sub>R PSC, respectively, one-way ANOVA). Furthermore, PPR with 50-ms interval was not significantly altered by increasing HEPES concentration from 0.8 to 10 mM in wild-type neurons (Fig. 6, C and D, n = 15, P = 0.81, paired t-test). This indicates that the difference in the PPR between wild-type and ASIC1 knockout neurons is not due to the low concentration of pH buffer (0.8 mM HEPES).

We also analyzed whether the reduction in the PPR was due to the loss of ASIC currents within the second EPSC. To test this, we measured residual currents in the presence of CNQX and AP5 in a paired-pulse protocol with a 50-ms interval (Fig. 6E). We did not observe a significant difference in the relative contribution of residual current to peak amplitude of AMPAR EPSC between the first and the second stimulation in either wild-type or ASIC1 knockout neurons (Fig. 6F, residual current after the first and the second stimulation = 4.7 ± 0.8 and 4.7 ± 0.6% of peak AMPAR EPSC, n = 5, P = 0.94 for wild-type; 5.5 ± 5.8 and 5.8 ± 0.78%, n = 4, P = 0.73 for ASIC1 knockout, paired t-test). Furthermore, inhibition of ASICs with amiloride did not affect the PPR with a 50-ms interval in either wild-type or ASIC1 knockout neurons (Fig. 6, G and H, 102 ± 2.6% of pre-amiloride control, n = 4, P = 0.51 for wild-type; 95 ± 2.7%, n = 4, P = 0.17 for ASIC1 knockout, paired t-test). These results suggest that the difference in PPR between wild-type and ASIC1 knockout neurons is not due to the direct contribution of ASIC current to AMPAR EPSC.

**Frequency of spontaneous neurotransmitter release is increased in ASIC1 knockout neurons**

Alterations in the PPR can be due to multiple factors. To further define the synaptic alterations in ASIC1 knockout neurons, we performed quantal analyses on spontaneous miniature (m) EPSCs of wild-type and ASIC1 knockout neurons in conventional mass culture (Mennerick et al. 1995). We recorded AMPAR-mediated mEPSC in extracellular solution containing 1 mM Mg<sup>2+</sup>, 30 μM bicuculline, and 1 μM tetrodotoxin to prevent action potential firing (Fig. 7A). The average peak amplitude of mEPSC was not different between wild-type and ASIC1 knockout neurons (Fig. 7B, 19.8 ± 2.2 pA, n = 11 neurons for wild-type, 21.1 ± 2.5 pA, n = 8 neurons for ASIC1 knockout, P = 0.70, unpaired t-test). In addition, neither the decay time constant nor the charge transfer of mEPSCs was different between wild-type and ASIC1 knockout neurons (decay time constant of mEPSC = 6.9 ± 0.1 and 7.1 ± 0.1 ms, for wild-type and ASIC1 knockout, P = 0.10, unpaired t-test; charge transfer of mEPSC = 102 ± 11 and 105 ± 12 fC for wild-type and ASIC1 knockout, P = 0.86, unpaired t-test). These results indicate that the postsynaptic response to spontaneous fusion of a single synaptic vesicle is not different in ASIC1 knockout neurons. However, mEPSCs were more frequent in ASIC1 knockout neurons than in wild-type neurons (Fig. 7C, 2.3 ± 0.2 Hz for wild-type; 5.1 ± 1.1 Hz for ASIC1 knockout, P < 0.05, unpaired t-test). An increase in mEPSC frequency is commonly caused by an increase in the total number of synapses, an increase in the size of the readily releasable pool (RRP) of synaptic vesicles, or an increase in the probability of neurotransmitter release. Previous studies have determined that the density of dendritic spines is equivalent in ASIC1 knockout and wild-type neurons in hippocampal slices (Zha et al. 2006). This suggests that the increased frequency of mEPSCs may be due to a larger number
FIG. 6. The effects of HEPES concentration and amiloride on AP-evoked postsynaptic currents and the PPR. A: representative traces of AMPAR EPSC, NMDAR EPSC, and GABA<sub>R</sub> PSC from wild-type neurons. Postsynaptic currents were recorded from the same autaptic neuron in 0.8 mM (black trace) and 10 mM (gray trace) HEPES-containing extracellular solutions. B: the effect of the HEPES concentration on postsynaptic currents. Postsynaptic currents were recorded in 0.8 mM and 10 mM HEPES-containing extracellular solutions from wild-type, ASIC1 knockout, or ASIC2 knockout neurons. Within each neuron, current amplitude in 10 mM HEPES solution was normalized to that in 0.8 mM HEPES solution. These relative current amplitudes (%) were not significantly different between wild-type, ASIC1 knockout, and ASIC2 knockout neurons (n = 15, P = 0.48, 0.83, and 0.60 for AMPAR EPSC, NMDAR EPSC, and GABA<sub>R</sub> PSC, respectively; one-way ANOVA). C: representative traces of AMPAR EPSC in a wild-type neuron evoked by the paired-pulse protocol of 50-ms interval in the extracellular solution containing 2 mM Ca<sup>2+</sup> and 1 mM Mg<sup>2+</sup> with 0.8 mM (black trace) and 10 mM (gray trace) HEPES. D: quantification of the effect of the HEPES concentration on the PPR. The PPR was not significantly different between 0.8 and 10 mM HEPES (n = 15, P = 0.81, paired t-test). E: representative traces of AMPAR EPSC and residual current in the presence of CNQX and AP5 in a wild-type autaptic neuron. AMPAR EPSC was recorded in the presence of AP5 (gray trace), and residual current was isolated using CNQX and AP5 (black trace). Voltage steps above the traces indicate transient membrane depolarization from −70 to 10 mV used to evoke APs with 50-ms interval. F: quantification of the ratio of the residual current in the presence of CNQX and AP5 to peak AMPAR EPSC in wild-type (n = 5) and ASIC1 knockout (n = 4) neurons. There is no significant difference in the contribution of residual current to peak AMPAR EPSC between the first and the second stimulation in either a wild-type or an ASIC1 knockout neuron (n = 5, P = 0.94 for wild-type; n = 4, P = 0.73 for ASIC1 knockout, paired t-test). G: representative traces of AMPAR EPSC evoked by the paired-pulse protocol of 50-ms interval before (−amiloride, black trace) and during 300 μM amiloride application (+amiloride, gray trace) in wild-type and ASIC1 knockout neurons. H: quantification of the effect of the amiloride on PPR. PPR during amiloride application (+amiloride) was normalized to PPR before amiloride treatment (−amiloride). Amiloride did not change PPR significantly in either wild-type or ASIC1 knockout neurons (n = 4, P = 0.51 for wild-type; n = 4, P = 0.17 for ASIC1 knockout, paired t-test).
of synaptic vesicles in the RRP or a higher probability of neurotransmitter release in ASIC1 knockout neurons.

**RRP size and refilling are not different in ASIC1 knockout neurons**

The RRP size and refilling were analyzed using hypertonic sucrose solution. When hypertonic sucrose is applied, synaptic vesicles in the entire RRP release neurotransmitters in a Ca\(^{2+}\)-independent manner and produce postsynaptic currents proportional to the size of the RRP (Rosenmund and Stevens 1996). Extracellular solution containing 500 mM sucrose was applied to neurons in mass culture for 4–5 s and the sucrose-induced postsynaptic current response was recorded in 1 mM Mg\(^{2+}\), 30 \(\mu\)M bicuculline, and 1 \(\mu\)M tetrodotoxin (Fig. 8A). This sucrose-induced response was completely inhibited by 10 \(\mu\)M CNQX (Fig. 8A), indicating that it was mediated by AMPARs (Rosenmund and Stevens 1996). RRP size was estimated from the sucrose-induced charge transfer calculated by integrating the transient component of the sucrose-evoked current. We determined that the RRP size was not significantly different between wild-type and ASIC1 knockout neurons (Fig. 8B, \(0.86 \pm 0.09\) nC, \(n = 20\) for wild-type, \(0.95 \pm 0.11\) nC, \(n = 24\) for ASIC1 knockout, \(P = 0.52\), unpaired \(t\)-test). The RRP refilling was also analyzed by comparing the charge transfer in response to a second sucrose application 3–4 s after the first (Fig. 8A) (Priller et al. 2006). RRP refilling, estimated as the ratio of the charge transfer by the second sucrose application to the charge transfer induced by the first sucrose application, was not significantly different between wild-type and ASIC1 knockout neurons (Fig. 8C, \(0.53 \pm 0.02\), \(n = 17\) for wild-type, \(0.52 \pm 0.03\), \(n = 20\) for ASIC1 knockout, \(P = 0.93\), unpaired \(t\)-test). These results indicate that the size and refilling of the RRP are not different between wild-type and ASIC1 knockout neurons in conventional mass culture.

**Progressive block of NMDARs by MK-801 was faster in ASIC1 knockout neurons**

An increase in the release probability could account for both the more frequent mEPSCs and the reduced PPR observed in ASIC1 knockout neurons (Zucker and Regehr 2002). The release probability of ASIC1 knockout neurons was analyzed using progressive block of the NMDARs by MK-801 (Rosenmund et al. 1993). Because MK-801 irreversibly blocks NMDARs while they are open, the rate of NMDAR EPSC decrease correlates with the probability of neurotransmitter release (Futai et al. 2007; Rosenmund et al. 1993). Thus the higher the release probability is, the faster the rate of MK-801-induced decrease of NMDAR EPSCs. The rate of progressive block of NMDAR by MK-801 could also be affected by the duration of NMDAR opening. Because NR2B-containing NMDARs open longer than NR2A-containing NMDARs, a difference in the ratio of NR2B- to NR2A-containing NMDARs can hamper interpretation of NMDAR progressive block by MK-801 (Monyer et al. 1992). However, we find that the weighted mean decay time constant of NMDAR EPSCs was not different between wild-type and ASIC1 knockout neurons (198 ± 14, \(n = 45\) for wild-type, and 182 ± 9, \(n = 48\) for ASIC1 knockout, \(P = 0.37\), unpaired \(t\)-test), suggesting that NMDARs remain open for similar periods of time in response to glutamate.

Under normal conditions, the NMDAR EPSCs of autaptic neurons stimulated every 10 s did not change significantly over time (Fig. 9A). In the presence of 10 \(\mu\)M MK-801, the NMDAR EPSCs gradually decreased in a stimulus-dependent manner (Fig. 9A). NMDAR EPSC amplitudes were normalized to the first NMDAR EPSC in the presence of MK-801 and plotted against stimulus number. The data were fit to a single-exponential equation, and the rate of the decrease of the NMDAR EPSCs was quantified by calculating the tau (\(\tau\)) in stimulus number (Fig. 9, B–D). The rate of NMDAR EPSC decrease in the presence of MK-801 was faster and the tau was significantly reduced in ASIC1 knockout neurons compared with wild-type neurons (Fig. 9, C and D, \(\tau = 22.4 \pm 1.9\) stimuli, \(n = 9\) for wild-type, 16.1 ± 1.5 stimuli, \(n = 8\) for ASIC1 knockout).
ASIC1 knockout, \( P < 0.05 \), unpaired \( t \)-test). These results suggest that the release probability is higher in ASIC1 knockout glutamatergic neurons.

**Neurotransmitter release probability is increased in ASIC1 knockout neurons**

A second experiment was used to assess the release probability of wild-type and ASIC knockout neurons. Both the AP-evoked EPSC and hypertonic sucrose-induced current were recorded in the same glutamatergic autaptic neurons (Fig. 10A). The probability of neurotransmitter release was calculated by dividing the charge transfer of AP-evoked EPSC by the charge transfer of the response induced by depletion of the RRP of synaptic vesicles with hypertonic sucrose (Molder et al. 2004; Rhee et al. 2002). The release probability of wild-type and ASIC2 knockout neurons was 6–7% (Fig. 10D, 7.0 ± 0.8%, \( n = 27 \) for wild-type, and 6.2 ± 0.5%, \( n = 12 \) for ASIC2 knockout), which is close to the value obtained in other studies (Rhee et al. 2002). The release probability of ASIC1 knockout neurons was significantly higher compared with wild-type and ASIC2 knockout neurons (Fig. 10D, 10.9 ± 1.7%, \( n = 19 \) for ASIC1 knockout, \( P < 0.05 \), ASIC1 knockout vs. wild-type or ASIC2 knockout, one-way ANOVA). Together, this experiment and the experiment assessing progressive block of NMDAR by MK-801 (Fig. 9) indicate that ASIC1 knockout neurons have a higher probability of neurotransmitter release than that of wild-type neurons.

**Rescue of ASIC1 knockout neurons by ASIC1a expression**

We find that the release probability is higher in neurons from ASIC1 knockout mice compared with neurons from wild-type mice. To determine whether this effect is the result of developmental compensation in response to ASIC1 gene disruption, we performed rescue experiments in cultured neurons. Hippocampal neurons from ASIC1 knockout mice were transfected during isolation with either vector alone or vector-expressing ASIC1a. Neurons were plated to foster microisland conditions, and whole cell patch clamping was used to measure acid-evoked currents and synaptic transmission after 14–20 days in culture. We found that extracellular acid (pH 6.0) failed to evoke \( H^+ \)-gated currents in ASIC1 knockout neurons transfected with vector alone (Fig. 11A, peak amplitude = 74 ± 17 pA, \( n = 9 \)). In ASIC1a-transfected ASIC1 knockout neurons, extracellular acid induced typical \( H^+ \)-gated currents with characteristics similar to those of wild-type neurons (Fig. 11A, peak amplitude = 1,489 ± 458 pA, \( n = 6 \), \( P < 0.05 \), ASIC1a-transfected vs. vector-transfected neurons, unpaired \( t \)-test). Thus transfection of ASIC1 knockout neurons with ASIC1a restored \( H^+ \)-gated currents. We next examined synaptic transmission in transfected ASIC1 knockout neurons. GABA\(_A\)-mediated postsynaptic currents were not significantly different in ASIC1 knockout neurons transfected with ASIC1a or vector alone (peak amplitude = 5.5 ± 1.3 nA, \( n = 6 \) for ASIC1a-transfected neurons, and 3.5 ± 0.8 nA, \( n = 8 \) for vector-transfected neurons, \( P = 0.23 \), unpaired \( t \)-test). However, the peak amplitude of AMPAR-mediated EPSCs was profoundly smaller in ASIC1a-transfected neurons compared with vector-transfected neurons (Fig. 11B, 1.60 ± 0.27 nA, \( n = 6 \) for ASIC1a-transfected neurons, and 4.42 ± 0.66 nA, \( n = 9 \) for vector-transfected neurons, \( P < 0.01 \), unpaired \( t \)-test). Furthermore, the PPRs at interpulse intervals of 40 and 50 ms were increased in ASIC1a-transfected neurons (Fig. 11C, \( P < 0.05 \), unpaired \( t \)-test). Because the PPR is inversely correlated with release probability (Zucker and Regehr 2002), these results suggest that release probability of ASIC1 knockout neurons
acidity is not surprising. For example, ASICs mediate acid-induced nociception during inflammation (Price et al. 2001; Sutherland et al. 2001). ASICs are involved in the retinal response to light where acidic pH changes are known to influence neuronal signaling (Ettaiche et al. 2004, 2006). In the brain, ASIC1a activation causes neuronal death during prolonged acidosis following ischemia (Xiong et al. 2004, 2006). However, ASICs are also involved in neuronal processes where the role of extracellular acidosis is not well defined. For example, ASIC1 is required for normal fear-related behaviors as well as learning and memory (Wemmie et al. 2002, 2003, 2004). Although rapid extracellular pH transients have been

transfected with ASIC1a is decreased. Moreover, we suggest that the decreased release probability also resulted in the smaller amplitude of AMPAR EPSCs in ASIC1a-transfected ASIC1 knockout neurons. These experiments show that the PPR of ASIC1 knockout neurons can be restored by in vitro expression of ASIC1a and suggest that the increase in release probability observed in ASIC1 knockout neurons is not due to developmental compensation.

**DISCUSSION**

Given that ASICs are activated by extracellular protons, a role for ASICs in biological processes involving extracellular
observed during synaptic transmission (Krishtal et al. 1987), how ASICs impact neurotransmission is not clear.

In this study, we investigated the role of ASICs in synaptic transmission by comparing postsynaptic currents of cultured hippocampal neurons from wild-type and both ASIC1 and ASIC2 knockout mice. We observed alterations in synaptic transmission in glutamatergic neurons from ASIC1 knockout mice using multiple experimental paradigms. First, the AMPAR/NMDAR EPSC ratio was reduced in ASIC1 knockout neurons (Fig. 2F). Second, the paired-pulse ratio (PPR) of AMPAR EPSC was reduced and the depression of AMPAR EPSCs during a short train of stimuli was greater in ASIC1 knockout neurons (Fig. 5). Alterations in the PPR can be due to both presynaptic and postsynaptic mechanisms. In an effort to identify the nature of the PPR changes in ASIC1 knockout neurons, we assessed other aspects of synaptic transmission. We determined that the quantal size of mEPSCs, the size of readily releasable pool (RRP) of synaptic vesicles, and RRP refilling were not significantly different in ASIC1 knockout neurons compared with wild-type neurons (Figs. 7 and 8). In contrast, the frequency of mEPSCs was increased (Fig. 7C), and the progressive block of NMDAR by MK-801 was faster in ASIC1 knockout neurons (Fig. 9). These results suggest that the reduced PPR in ASIC1 knockout neurons is due to the increased probability of neurotransmitter release. This was confirmed by the observation that the release probability, as measured by the ratio of AP-evoked to hypertonic sucrose-evoked charge transfer, was increased in ASIC1 knockout neurons (Fig. 10). Together, these results support the conclusion that cultured ASIC1 knockout neurons have a higher release probability than that of wild-type neurons. Further, the mEPSC frequency increased and the release probability was augmented without a change in the RRP size suggest an enhanced fusion propensity of individual vesicles in ASIC1 knockout neurons. Transfection of ASIC1a increased the PPR of ASIC1 knockout neurons and reduced AMPAR EPSCs (Fig. 11). These results are consistent with an ASIC1a-dependent rescue of the release probability of ASIC1 knockout neurons and suggest that the increased release probability observed in ASIC1 knockout neurons is not likely due to developmental compensation. Interestingly, there were no significant differences in synaptic transmission in ASIC2 knockout neurons (Figs. 2F, 5B, and 10D), indicating that these alterations are specific to disruption of ASIC1.

Because the release probability is increased in ASIC1 knockout neurons, AP-evoked whole cell postsynaptic currents should be larger compared with wild-type neurons. Although the average peak amplitudes of single AP-evoked AMPAR EPSCs or NMDAR EPSCs were larger in ASIC1 knockout neurons (Figs. 2, D and E and 10B), the difference did not reach statistical significance, and we expect a greater increase in EPSC amplitude with such an increase in release probability.

![Figure 11](http://jn.physiology.org/) Rescue of ASIC1 knockout neurons by ASIC1a expression. A: representative traces of H⁺-gated currents in ASIC1 knockout neurons transfected with vector (pcDNA3.1, left) or human ASIC1a (right). To evoke H⁺-gated currents, we exchanged the extracellular solution of pH 7.4 to pH 6.0 for 6–7 s (gray bars). B: peak amplitude of pH 6.0–induced H⁺-gated currents. n = 6 for ASIC1a-transfected neurons; n = 9 for vector-transfected neurons; unpaired t-test. C: representative traces of whole cell AMPAR-mediated EPSC in ASIC1 knockout neurons transfected with vector or ASIC1a. AMPAR EPSCs were recorded in the presence of 1 mM Mg²⁺, and membrane potential was held at −70 mV. D: peak amplitude of the AMPAR EPSCs. n = 6 for ASIC1a-transfected neurons; n = 9 for vector-transfected neurons; unpaired t-test. E: representative traces of AMPAR EPSCs evoked by a paired pulse with 50-ms interval in the presence of 2 mM Ca²⁺ and 1 mM Mg²⁺. F: quantification of the PPR at intervals from 20 to 50 ms. The PPR was calculated as the ratio of the peak amplitude of the second AMPAR EPSC to the peak amplitude of the first AMPAR EPSC. n = 4–6 for ASIC1a-transfected neurons; n = 5–9 for vector-transfected neurons. *P < 0.05, unpaired t-test. Single glutamatergic neurons on microisland were selected for recording H⁺-gated currents and AMPAR EPSC in these experiments.
We suggest that the postsynaptic response of ASIC1 knockout neurons is altered such that the EPSC amplitude is unchanged even with the increased release probability. In support of this idea, we did observe that AMPAR/NMDAR EPSC ratio was reduced in ASIC1 knockout neurons (Fig. 2F). Because the quantal size of AMPAR-mediated mEPSC was not changed in ASIC1 knockout neurons (Fig. 7B), it is not likely that the number of AMPARs per synapse or the conductance of individual postsynaptic AMPARs was different in the ASIC1 knockout neurons. In contrast, disruption of ASIC1 may have affected the ratio of AMPAR-expressing functional synapses to silent synapses that contain NMDAR but not AMPAR. Thus the number of AMPAR-expressing functional synapses may be reduced in ASIC1 knockout neurons. Such a reduction would mask the effect of enhanced release probability on AMPAR EPSCs. In rescue experiments, we did observe alterations in postsynaptic current consistent with altered release probability. ASIC1 knockout neurons transfected with ASIC1a displayed dramatically reduced AMPAR EPSCs compared with vector-transfected neurons, consistent with a decrease in release probability (Fig. 11B). This result suggests that presynaptic alterations in ASIC1 knockout neurons can be rescued by expression of ASIC1a in culture.

It is hypothesized that postsynaptic ASICs are activated by protons released from synaptic vesicles during neurotransmission. ASIC activation, in turn, contributes to depolarization and calcium-induced signaling cascades. This model has been difficult to prove and, like others before, we did not detect ASIC-mediated postsynaptic currents during synaptic transmission (Fig. 3, A–D). Even multiple stimulations failed to induce measurable ASIC-dependent currents (Fig. 6, E and F). Furthermore, we did not observe any ASIC-dependent effect of amiloride on AMPAR EPSC or PPR (Figs. 3, E and F and 6, G and H). Given the large postsynaptic currents recorded, the sensitivity of the method, and the presence of H+–gated currents evoked by exogenous acid application in these neurons, we conclude that specific conditions must exist for ASICs to be activated in this manner and make a substantial contribution to the postsynaptic currents. However, definitive changes in synaptic transmission were observed in ASIC1 knockout neurons, even though ASIC currents were not detected during synaptic transmission. Our data suggest that the higher release probability in ASIC1 knockout neurons is due to either 1) disruption of ASIC1a-specific signaling, which has long-term effects on presynaptic mechanisms, or 2) loss of an unconventional nonionotropic ASIC function, which is not dependent on ion conduction of ASIC1a.

First, disruption of ASIC1a-specific signaling could have long-term effects on presynaptic mechanisms. Under normal conditions, ASIC1a activation could lead to long-term changes in synaptic transmission through activation of signal transduction cascades. ASIC1a homomultimers are Ca2+-permeable (Yermolaieva et al. 2004) and may activate Ca2+/calmodulin-dependent protein kinase II (CaMKII). Phosphorylated CaMKIIα is reduced in brains from ASIC1 knockout mice (Zha et al. 2006), indicating that disruption of ASIC1 can influence the activation status of CaMKII, which is essential for the recruitment of AMPAR to synapses and for the LTP induction in the CA1 region of hippocampus (Lisman et al. 2002). CaMKII may also affect presynaptic mechanisms to alter neurotransmitter release (Chi et al. 2001; Llinàs et al. 1991; Sanhueza et al. 2007; Waxham et al. 1993). To investigate the possibility that previous activation of ASIC1a causes changes in synaptic transmission, we assessed postsynaptic current following activation of ASICs by exogenous acid application. However, we did not observe changes in postsynaptic currents in the time frame assessed (2–5 min, Fig. 4). Thus ASIC1a-induced changes in signal transduction may occur under other specific conditions or require a longer period of time to manifest.

Second, changes in synaptic transmission of ASIC1 knockout neurons could be due to the loss of an unconventional nonionotropic ASIC function. Such “nonconducting” functions have been reported for other ion channels such as NMDARs (Alvarez et al. 2007) and HCN (Ih) channels (Beamont and Zucker 2000; Beamont et al. 2002; Zhong et al. 2004). ASIC1a could regulate release probability by direct interaction with neurotransmitter release machinery. Previous studies indicate that ASICs are localized postsynaptically (Wemmie et al. 2002; Zha et al. 2006), but do not exclude the possibility of ASIC1a at presynaptic sites (Alvarez de la Rosa et al. 2003). Ionotropic neurotransmitter receptors such as NMDA receptor, kainite receptor, and GABA_A receptor are expressed both postsynaptically and postsynaptically and regulate neurotransmitter release (MacDermott et al. 1999). Furthermore, ASIC1a, ASIC2a, and γ-ENaC form heteromeric channels in malignant glioma cells, and this complex interacts with syntaxin 1A, a component of the SNARE complex involved in exocytosis of synaptic vesicles (Berdiev et al. 2003). Although it is not known whether this interaction also occurs in neurons, this raises a possibility that ASICs might directly regulate neurotransmitter release. Alternatively, postsynaptic ASIC1a could affect synaptic transmission in a retrograde manner like postsynaptic PSD-95, which regulates neurotransmitter release by physical interaction with presynaptic neurexin (Futaí et al. 2007). ASICs have a large extracellular loop of largely unknown function, and the extracellular loop of postsynaptic ASIC1a could physically interact with a presynaptic counterpart to directly regulate neurotransmitter release.

ASIC1 knockout mice exhibited defects in multiple aspects of learning and memory such as spatial learning, fear conditioning, and eye-blink conditioning (Wemmie et al. 2002, 2003). This suggests a fundamental role of ASIC1a in synaptic transmission and plasticity. Our results indicate that disruption of ASIC1 increases presynaptic neurotransmitter release. In the brain, the greater basal release probability might prevent additional increases in synaptic transmission and thus limit long-term potentiation in ASIC1 knockout mice (Wemmie et al. 2002). We observed altered paired-pulse modulation and depression during short trains of stimulation in ASIC1 knockout neurons. These types of short-term plasticity are important for information processing in neural networks (e.g., producing reliable response to repetitive activation) and contribute to normal behavior (Blitz et al. 2004). Thus dysregulation of short-term plasticity may also cause defects in multiple aspects of cognitive function in ASIC1 knockout mice. Our results indicate that the consequences of ASIC1 disruption are complex and influence both presynaptic and postsynaptic mechanisms. In addition, ASIC1a may play a fundamental role in synaptic transmission by regulating presynaptic release probability.
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