Cellular/Molecular

Differential Responses of Bladder Lumbosacral and Thoracolumbar Dorsal Root Ganglion Neurons to Purinergic Agonists, Protons, and Capsaicin

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The present study explored differences in sensitivity to purinergic agonists, protons, and capsaicin in lumbosacral (LS) and thoracolumbar (TL) sensory neurons that innervate the rat urinary bladder. The majority of LS neurons (93%) were sensitive to α,β-methyleneATP (α,β-metATP) compared with 50% of TL neurons. Based on inactivation kinetics, a slowly desensitizing current evoked by α,β-metATP predominated in LS neurons (86%) compared with mixed components that characterized TL neuron responses (58%). The density of the slowly desensitizing current was greater in LS than in TL neurons (LS, 34.4 ± 5.3 pA/pF; TL, 2.5 ± 0.8 pA/pF). Almost all neurons in both ganglia responded to protons and to capsaicin (LS, 100%; TL, 98%). Proton-activated currents in bladder sensory neurons exhibited distinct inactivation kinetics as fast, intermediate, slowly desensitizing, and sustained components. More than one component was expressed in every cell. Although there was no difference in the percentage of neurons expressing more than one component, the density of the sustained current was significantly greater in LS than in TL neurons (LS, 86.1 ± 7.6 pA/pF; TL, 30.3 ± 7.0 pA/pF). Similarly, the capsaicin-evoked current was greater in LS than in TL neurons (LS, 129.6 ± 17.4 pA/pF; TL, 86.9 ± 11.6 pA/pF). Finally, a greater percentage of TL neurons bound isolectin B4 than LS neurons (LS, 61%; TL, 85%). The greater degree of α,β-metATP, proton, and capsaicin responsiveness, in addition to differences in current type and current densities, in LS and TL neurons suggests that bladder pelvic and hypogastric/lumbar splanchnic afferents are functionally distinct and likely mediate different sensations arising from the urinary bladder.

Key words: bladder; dorsal root ganglia; α,β-metATP; ATP; P2X receptors; protons; capsaicin; IB4; A317491

Introduction
As a temporary storage organ, the urinary bladder is exposed to varying volumes of urine of differing composition, which ultimately triggers micturition and also sensation. Importantly, interactions between the bladder urothelium and terminals of sensory neurons, which can be activated by mechanical and/or chemical stimuli, play a significant role in both the regulation of micturition and sensations arising from the bladder (Andersson, 2002). For example, distension of the bladder releases ATP from urothelial cells, which activates sensory neurons, potentially triggering reflex micturition (Vlaskovska et al., 2001). Deletion of P2X3 receptors in mice results in bladder hyporeflexia, increased voiding volume, and decreased micturition frequency (Cockayne et al., 2002). Similarly, activation of the capsaicin receptor TRPV1 contributes to the micturition reflex, as shown by bladder hyporeflexia in TRPV1 knock-out mice and an association between detrusor hyperactivity and increased TRPV1 expression (Birder et al., 2002; Brady et al., 2004).

The bladder is innervated by the pelvic and hypogastric/lumbar splanchnic nerves with cell bodies located in thoracolumbar (TL) and lumbosacral (LS) dorsal root ganglia (DRGs), respectively (Applebaum et al., 1980). Most studies have focused on the pelvic nerve, transection of which abolishes bladder contractions to repetitive filling, suggesting that the pelvic nerve is important for the sensation of bladder distension and micturition (Kontani and Hayashi, 1997; Meen et al., 2001). However, the hypogastric/lumbar splanchnic nerve is also activated by mechanical and chemical stimulation of the bladder, contributes to the regulation of micturition, and may play an enhanced role in cystitis (Moss et al., 1997; Mitsu et al., 2001).

Accumulating evidence suggests that different nerves innervating the same organ are distinct functionally. For example, the colon is also innervated by the pelvic and hypogastric/lumbar splanchnic nerves, and the mechanosensitivity and location of receptive endings in the colon differ between the two nerves (Brierley et al., 2004), as do the passive and active electrical properties of colonic TL and LS neurons (Gold and Traub, 2004). In addition, hypogastric/splanchnic sensory afferents reportedly respond more robustly to chemical stimuli than do pelvic counterparts (Moss et al., 1997). Finally, the innervations of the bladder

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body and proximal urethra tract exhibit distinct histological and electrical properties in addition to differential expression of TRPV1 and voltage-gated Na⁺ and Ca²⁺ currents (Yoshimura et al., 2003). Despite such evidence suggesting that bladder sensory afferent pathways are distinct functionally, there has been no systematic study of how bladder LS and TL neurons differ, which could further support/advance our knowledge of micturition and sensory function of the bladder in normal and pathological states.

Considering the importance of P2X, TRPV1, and, potentially, acid-sensing ion channels (ASICs) in micturition and bladder sensory function, we hypothesized that the properties of urinary bladder sensory neurons differ in TL and LS DRGs and examined in the present study purinergic, proton, and capsaicin sensitivity of retrogradely labeled bladder neurons. A preliminary report of these results has been published previously in abstract form (Dang et al., 2004).

Materials and Methods

Male Sprague Dawley rats (200–300 g; Harlan, Indianapolis, IN) were used for the experiments. Rats were housed under a 12 h light/dark cycle with ad libitum access to food and water. Animal handling adhered to the National Research Council Guide for the Care and Use of Laboratory Animals; the experimental protocol was approved by the Animal Care and Use Committee of The University of Iowa.

Cell labeling

Under pentobarbital anesthesia (50 mg/kg, i.p.), the rat bladder was exposed surgically (lower abdominal incision, ~1 cm in length), and 1,1'-dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine methanesulfonate (DiI; 100 mg in 2 ml of DMSO; Molecular Probes, Eugene, OR) was injected into six to eight sites within the wall of the bladder base around the trigone using a 30 gauge needle (6 µl per site). Any visible leakage of DiI during injection was wiped off with a cotton swab. The incision was closed with a 4.0 silk suture, and rats were allowed to recover until the DRGs were harvested 14–21 d later for electrophysiological recordings. We checked every rat postmortem for leakage of DiI by examining the bladder and surrounding tissues after the TL or LS DRGs were removed. No labeling in any tissue other than the bladder was noted.

Cell dissociation and culture

Rats were anesthetized as above and decapitated, and the T13–L2 or L6–S2 DRGs were removed rapidly. The ganglia were minced and incubated at 37°C in 5% CO₂ for 60 min in serum-free, supplemented Neuro-A medium (5% B27 supplements, 0.5 mM L-glutamine, and 1% glucose). The pH was adjusted to 7.4 with NaOH (310 mOsm). Neu- rons that innervated the bladder were identified by DiI content using a rhodamine filter (excitation wavelength, ~546 nm; barrier filter, 580 nm). Fire-polished micropipettes with tip resistances of 1.5–2 MΩ were used for current- and voltage-clamp recordings. The uncompensated series resistance was generally approximately ≤7 MΩ. The pipette was filled with an internal solution consisting of the following (in mM): 130 KC1, 1 CaCl₂, 1 MgCl₂, 10 EGTA, 10 HEPES, 4 Na₃ATP, 0.5 Tris-GTP, and 0.5 GDP. The pH was adjusted to 7.2 using KOH (310 mOsm). After establishing the whole-cell configuration, the voltage was clamped at ~70 mV using an Axopatch 200B amplifier (Axon Instruments, Union City, CA), digitized at 1 kHz (Digidata 1350; Axon Instruments), and controlled using Clampex software (pClamp 9; Axon Instruments). Cell capacitance was obtained by reading the value from the Axopatch 200B amplifier. Recordings began 2–3 min after establishing the whole-cell configuration to ensure stable recording conditions.

Drugs were applied using a fast-step SF-77B superfusion system (Warner Instruments, Hamden, CT) with a three-barrel pipette placed in close proximity (100 µm) to the cell. Complete solution exchange could be obtained within 24 ± 1 ms by measuring the time constant from switching from the standard external solution to one containing 40 mM KC1. Antagonists were applied for 4 s, whereas agonists were superfused for 30 s before the application of agonists. A washout period of 4 min was allowed between agonist applications. Drugs were obtained from Sigma-Aldrich (St. Louis, MO) and prepared fresh from stock solutions on the day of the experiment. All experiments were performed at room temperature (21–23°C).

Immunohistochemistry

Isolecitin B4 labeling. After the recording protocol was completed, isolecitin B4 (IB4) conjugated to AlexaFluor 488 (Molecular Probes, Eugene, OR) was applied to the recording chamber for 10 min at a final concentration of 10 µg/ml (from a stock concentration of 1 mg/ml). The recording chamber was then washed for 3 min to remove any excess conjugated IB4. IB4-positive neurons displayed a bright green ring around the cell surface when barrier filtered at 510 nm. To confirm the specificity of IB4 labeling, DRG cells were preincubated with excess unconjugated IB4 (0.2 mg/ml; Sigma-Aldrich) before applying conjugated IB4. No labeling was observed (data not shown). In addition, we also compared IB4 labeling using FITC–IB4 (Sigma-Aldrich) and AlexaFluor 488–IB4 at the same concentration (2 µg/ml) in serially cut DRG slices. AlexaFluor 488–IB4 always labeled more neurons than FITC–IB4 (data not shown); however, we did not quantify the percentages of IB4-positive neurons labeled with FITC or AlexaFluor 488 in DRG sections.

P2X and TRPV1 labeling. To examine colocalization of receptors, we performed triple labeling of P2X2 and P2X3, or P2X2, or TRPV1, or TRPV1. Fluorogold (4%; Biotium, Hayward, CA) was injected into the bladder base in three rats as described for DiI above. Rats were killed 14 d later, and DRGs were rapidly removed, fixed in 4% paraformaldehyde (Sigma-Aldrich) for 4 h, and cryoprotected in 30% sucrose for 3 d before cutting. DRGs were sectioned at 10 µm with a Leica (Nussloch, Germany) 3M 3050 cryostat. DRG sections were blocked with 3% goat serum in 0.01M PBS (Sigma-Aldrich) for 2 h before incubation of primary antibodies for 24 h at 4°C. Primary antibodies (rabbit anti-rat) were applied at the following dilutions: P2X2, 1:100; P2X3, 1: 200; TRPV1, 1:50 (EMD Biosciences, La Jolla, CA). Primary antibodies were aspirated, and sections were washed three times with 0.01M PBS. Secondary antibodies (anti-rabbit) conjugated to either AlexaFluor 488 or 568 (Molecular Probes) were applied for 4 h and washed three times with 0.01M PBS. Sections were mounted with fluorescence mount (Sigma-Aldrich) and viewed with a Nikon microscope equipped with separate fluorescence filters. Images were captured with a model 2.3.1 SPOT digital camera (Diagnostic Instruments, Sterling Heights, MI). For control experiments, primary antibodies were omitted and/or blocked with corresponding antigenic peptides for 24 h. No labeling was observed (data not shown).

Data analyses

All data are given as mean ± SEM. Analyses were performed using the software program Graph Pad Prism 4 (Graph Pad Software. San Diego, CA). Sigmoidal dose–response curves were generated using the following equation: \( Y = A/[1 + \exp\left(-\log(E_{50} - X)/B\right)] \), where \( A \) is the loga- rithm of concentration and \( Y \) is the response and starts at 0 and goes to \( A \) with a sigmoidal shape. \( B \) is the Hill slope. Desensitization kinetics were fitted with a standard exponential equation: \( Y = K_0 + K_1 \times \exp(-t/r) \),
where $Y$ is the current amplitude at time $t$, $K_0$ is the amplitude of the sustained component, and $\tau$ is the time constant. $K_0$ and $K_1$ represent the contribution to current amplitude from the fast and slow components of the current, respectively.

To compare percentage differences, a $\chi^2$ test was used. All other comparisons were made using Student’s unpaired $t$ test. Results were considered to be statistically significant when $p < 0.05$.

**Results**

**Bladder innervation density and cell size characteristics**

A total of six rats were used to estimate the number of primary afferent neurons that innervate the bladder. We randomly chose six coverslips (one from each rat) containing T13–L2 or L6–S2 DRGs and counted the total number of DiI-labeled neurons. In the L6–S2 DRGs, 219 of 4015 neurons were DiI labeled, constituting $5.4 \pm 0.5\%$ of the sample. In contrast, only $2.0 \pm 0.3\%$ (106 of 5316) of T13–L2 DRG neurons were DiI positive ($p < 0.05$), suggesting that the number of bladder DRG neurons is ~2.5-fold greater in LS than in TL DRGs.

Next, we examined whether whole-cell capacitance (as an index of cell size) differed between LS and TL bladder neurons. As illustrated in Figure 1A, the distribution of cell capacitances in LS and TL bladder neurons differed significantly. A significant portion of LS bladder neurons (49 of 194; 25.3%) had a whole-cell capacitance <20 pF, suggesting that these neurons were small (Fig. 1B). Most remaining neurons (144 of 194; 74%) had capacitances between 20 and 60 pF, consistent with medium size. In contrast, no TL bladder neurons had a whole-cell capacitance <20 pF ($p < 0.001$ vs LS bladder neurons), whereas 19 of 123 (15.4%) were considered to be large because cell capacitance was >60 pF ($p < 0.01$); most (104 of 123; 85.4%) TL bladder neurons were medium-sized cells (Fig. 1B). Consistent with the difference in size distribution, the mean capacitance of LS bladder neurons (27.7 ± 0.7 pF) differed significantly from the mean capacitance of TL bladder neurons (45.3 ± 1.3 pF; $p < 0.001$).

**Characterization of bladder sensory neuron responses to ATP**

**LS bladder neurons**

To evaluate functionally the presence of purinergic receptors on bladder neurons, we applied the nonselective agonist ATP. ATP superfusion triggered rapidly activating inward currents in 53 of 54 (98.1%) LS neurons (Fig. 2A–C), consistent with the presence of P2X receptors. Based on the kinetics of current decay, we differentiated fast, slow, and mixed desensitizing currents, with a predominance of the slowly desensitizing current found in 46 of 53 (86.8%) neurons (Fig. 2B). Three (5.7%) and four (7.5%) LS neurons expressed fast (Fig. 2A) and mixed (Fig. 2C) current kinetics, respectively. The fast desensitizing current peaked within 237.5 ± 67 ms and desensitized with a time constant of 241.2 ± 42 ms. The slowly desensitizing current reached the maximal amplitude within 1.03 ± 0.07 s and decayed with a time constant of 4.44 ± 0.42 s. The sample trace shown in Figure 2C suggests that the slowly desensitizing component contributed significantly to the response in cells with mixed current kinetics. Consistent with this visual impression, a double-exponential fit revealed that the slow component amounted to 83 ± 6% of the peak current. The mean current density for fast, mixed, and slowly desensitizing components was 5.3 ± 0.6, 9.9 ± 2.5, and 75.3 ± 14 pA/pF, respectively.

**TL bladder neurons**

In contrast to LS neurons, application of ATP evoked currents in only 34 of 52 (65.4%; $\chi^2 = 17.2; p < 0.01$) TL neurons. Thirteen of 34 (38.2%), 4 of 34 (11.8%), and 17 of 34 (50%) TL neurons expressed slowly, fast, and mixed desensitizing currents, respectively (Fig. 2D–F). The slowly desensitizing component peaked within 1445 ± 168 ms and decayed with a time constant of 3.75 ± 0.9 s. The fast component peaked within 295.7 ± 26 ms and had a desensitization constant of 309.8 ± 24 ms. The mixed current peaked just as rapidly as the fast desensitizing component, with a time-to-peak of 292.5 ± 21 ms. Based on exponential fitting, the slowly desensitizing current accounted for 42 ± 7% of the peak current. The mean current density for fast, mixed, and slowly desensitizing currents was 4.3 ± 0.4, 7.3 ± 1.5, and 4.6 ± 0.7 pA/pF, respectively. Although the current density of fast and mixed currents did not differ between LS and TL neurons, LS neurons with slowly desensitizing kinetics expressed significantly greater peak current densities ($p < 0.001$). Similarly, in LS neurons with mixed kinetics, the slow component contributed a greater
percentage of the peak current compared with TL counterparts \((p < 0.001)\). Differences between LS and TL neurons are summarized in Table 1.

Characterization of P2X receptor-mediated currents in bladder sensory neurons

**LS bladder neurons**

Because ATP is a nonselective purinergic agonist, we examined the effects of \(\alpha\beta\)-methyleneATP (\(\alpha\beta\)-metATP), a selective homomeric P2X1, homomeric P2X3, and heteromeric P2X2/3 receptor agonist (Dunn et al., 2001). Application of \(\alpha\beta\)-metATP evoked an inward current in 50 of 54 (93%) LS bladder neurons (Fig. 2A–C). In three neurons, only ATP and not \(\alpha\beta\)-metATP triggered inward currents. Based on the current kinetics described above, we identified a slow current in 43 of 50 (86%) (Fig. 2B), a fast current in 3 of 50 (6%), and a mixed current in the remaining 4 of 50 (8%) LS neurons. The activating and desensitization kinetics were similar to those obtained in response to ATP and are summarized in Table 2. In neurons with mixed kinetics, the slowly desensitizing component did not decay sufficiently for analysis, and we therefore could not estimate the contribution of the slowly desensitizing current to the total current. The mean fast, mixed, and slowly desensitizing currents in LS neurons were 2.5 ± 0.8, 4.2 ± 1.0, and 34.4 ± 5.3 pA/pF, respectively, with neurons expressing the slowly desensitizing component having the greatest peak current density \((p < 0.001)\). Compared with the non-specific agonist ATP, peak current densities were significantly less with \(\alpha\beta\)-metATP \((p < 0.05)\).

**TL bladder neurons**

As was seen for LS bladder neuron responses to ATP and \(\alpha\beta\)-metATP, \(\alpha\beta\)-metATP did not trigger inward currents in all TL bladder neurons that responded to ATP. Only 26 of 52 (50%) TL neurons responded to \(\alpha\beta\)-metATP (comparison between LS and TL neurons, \(\chi^2 = 21.6; p < 0.01\)). In eight TL neurons, ATP but not the more selective P2X agonist \(\alpha\beta\)-metATP triggered inward currents that slowly activated and desensitized. Similar to the effects of ATP, \(\alpha\beta\)-metATP triggered slow currents in 5 of 26 (19.2%), fast currents in 6 of 26 (23.1%), and mixed currents in 15 of 26 (57.7%) TL neurons. Current kinetics are summarized in Table 2. Based on exponential fitting, the slowly desensitizing component contributed 29.4 ± 5% of the peak current in cells with mixed kinetics. When normalized to cell size, the mean current density for fast, mixed, and slowly desensitizing currents was 1.8 ± 0.3, 2.6 ± 0.3, and 2.5 ± 0.8 pA/pF, respectively. Although there was no difference between densities of \(\alpha\beta\)-metATP-evoked currents in TL neurons, the slowly desensitizing current density was significantly less in TL than in LS neurons \((p < 0.001)\). In addition, peak current densities were significantly less after activation with \(\alpha\beta\)-metATP when compared with the non-specific agonist ATP \((p < 0.05)\).

### Table 1. Summary of frequency of responses to and current densities evoked by ATP, \(\alpha\beta\)-metATP, protons, and capsaicin in bladder sensory neurons

<table>
<thead>
<tr>
<th></th>
<th>LS neurons</th>
<th>TL neurons</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ATP</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage responding</td>
<td>98% (53/54)</td>
<td>65% (34/52)*</td>
</tr>
<tr>
<td>(I_d) density (pA/pF)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>5.3 ± 0.6</td>
<td>4.3 ± 0.4</td>
</tr>
<tr>
<td>Slow</td>
<td>75.3 ± 14</td>
<td>4.6 ± 0.7*</td>
</tr>
<tr>
<td>Mixed</td>
<td>9.9 ± 2.5</td>
<td>7.3 ± 1.5</td>
</tr>
<tr>
<td><strong>(\alpha\beta)-metATP</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage responding</td>
<td>93% (50/54)</td>
<td>50% (26/52)*</td>
</tr>
<tr>
<td>(I_d) density (pA/pF)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>2.5 ± 0.8</td>
<td>1.8 ± 0.3</td>
</tr>
<tr>
<td>Slow</td>
<td>34.4 ± 5.3</td>
<td>2.5 ± 0.8*</td>
</tr>
<tr>
<td>Mixed</td>
<td>4.2 ± 1</td>
<td>2.6 ± 0.3</td>
</tr>
<tr>
<td><strong>Protons</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage responding</td>
<td>78% (42/54)</td>
<td>87% (45/52)</td>
</tr>
<tr>
<td>(I_d) density (pA/pF)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>56.4 ± 10</td>
<td>48.2 ± 10</td>
</tr>
<tr>
<td>Intermediate</td>
<td>57.8 ± 11</td>
<td>59 ± 12</td>
</tr>
<tr>
<td>Slow</td>
<td>52.2 ± 16</td>
<td>74.8 ± 15</td>
</tr>
<tr>
<td>Sustained</td>
<td>86.1 ± 16</td>
<td>30.3 ± 7*</td>
</tr>
<tr>
<td><strong>Capsaicin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Percentage responding</td>
<td>91% (49/54)</td>
<td>92% (48/52)</td>
</tr>
<tr>
<td>(I_d) density (pA/pF)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>130 ± 17</td>
<td>87 ± 11*</td>
</tr>
</tbody>
</table>

Data are a comparison between LS and TL neurons \((p < 0.05)\).

**P2X receptor dose–response relationship**

We next determined the dose–response relationship for the two purinergic agonists by applying either \(\alpha\beta\)-metATP or ATP in increasing concentrations. To control for differences in current densities, we expressed current as a fraction of relative current at...
Table 2. Summary of purinergic agonist-evoked current kinetics in bladder sensory neurons

<table>
<thead>
<tr>
<th></th>
<th>ATP</th>
<th>αβ-metATP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ttp (ms)</td>
<td>tau 1 (ms)</td>
</tr>
<tr>
<td>LS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>238 ± 67</td>
<td>241 ± 42</td>
</tr>
<tr>
<td>Slow</td>
<td>1030 ± 70</td>
<td>4440 ± 420</td>
</tr>
<tr>
<td>Mixed</td>
<td>255 ± 50</td>
<td>241 ± 18</td>
</tr>
<tr>
<td>TL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>296 ± 26</td>
<td>310 ± 24</td>
</tr>
<tr>
<td>Slow</td>
<td>1445 ± 168</td>
<td>3750 ± 900</td>
</tr>
<tr>
<td>Mixed</td>
<td>293 ± 21</td>
<td>239 ± 26</td>
</tr>
</tbody>
</table>

ttp, Time to peak; n/a, not applicable.

Figure 3. A, B, Dose–response relationship of slowly (A) and rapidly (B) desensitizing currents in response to application of αβ-metATP or ATP. ATP (filled circles) was more potent than αβ-metATP (open circles) with EC50 values of 44.6 ± 4 μM (n = 17) and 17.2 ± 3 μM (n = 11), respectively, for the slowly desensitizing current. Similarly, ATP was more potent than αβ-metATP with EC50 values of 36.6 ± 5 μM (n = 3) and 20.2 ± 4 μM (n = 4), respectively, for the fast desensitizing current. C, D, Recovery kinetics for the slowly (C) and rapidly (D) desensitizing currents by varying the interpulse intervals of αβ-metATP (30 μM). E, Recovered current fractions were fitted with a single exponential. Although the slow current (open circle) minimally desensitized and recovered rapidly with a constant of 10.8 ± 5.7 s (n = 3), the fast desensitizing current (filled circle) recovered significantly slower with a constant of 45 ± 3 s (n = 3; p < 0.001).

300 μM. As shown in Figure 3, A and B, both αβ-metATP- and ATP-activated currents were concentration dependent for both the slowly desensitizing and rapidly (fast) desensitizing currents. The EC50 values for αβ-metATP and ATP for the slowly desensitizing current were 44.6 ± 4 μM (n = 17) and 17.2 ± 3 μM (n = 11), respectively (p < 0.001). The EC50 values for αβ-metATP and ATP for the rapidly desensitizing current were 36.6 ± 5 μM (n = 3) and 20.2 ± 4 μM (n = 4), respectively (p < 0.05). ATP was found to be significantly more potent than αβ-metATP for activation of the two kinetically distinct currents. To further examine whether ATP activates other P2 receptors, we applied maximal concentrations of αβ-metATP (300 μM) or ATP (300 μM) to some neurons. ATP produced a 43.4 ± 5.5% (n = 6; p < 0.001) greater peak current than did αβ-metATP for the slowly desensitizing current. In contrast, αβ-metATP and ATP evoked similar peak current for the fast desensitizing current (4.7 ± 1.2%; n = 4; p > 0.05). Therefore, the greater potency of ATP relative to αβ-metATP, and activation of other P2 receptors, appears to contribute to the greater slowly desensitizing current observed with ATP. However, the greater potency of ATP relative to αβ-metATP accounts for the larger fast desensitizing current described after administration of ATP.

**Recovery kinetics**

To further characterize the αβ-metATP-evoked responses, we examined the recovery kinetics from desensitization of αβ-metATP-elicited currents in principal current types found in LS neurons (slowly desensitizing) and TL neurons (fast/mixed desensitizing currents). The application duration remained the same (4 s) in these experiments, but the application interval was increasingly extended, and the fraction of current in response to the second application of αβ-metATP relative to the first application was plotted against time using a single-exponential fit. The slowly desensitizing current showed little decay and recovered rapidly (Fig. 3C), whereas the fast desensitizing component required 3 min for complete recovery (Fig. 3D).

Based on single-exponential fitting (Fig. 3E), the slowly desensitizing current recovered significantly faster than the rapidly desensitizing current (time constants of 10.8 ± 5.7 s (n = 3) and 45.5 ± 3 s (n = 3), respectively; p < 0.001). Because rapidly and slowly desensitizing currents were rarely seen in LS and TL bladder neurons, respectively, we did not systematically examine their recovery kinetics.

**Antagonism of αβ-metATP-evoked currents by selective antagonists**

To facilitate the construction of dose–inhibition relationships for antagonists, we examined their effects using LS neurons because they showed a higher frequency of response than TL neurons. Once IC50 values were determined, excess concentrations of antagonist were used to examine αβ-metATP-evoked currents. To confirm that αβ-metATP-evoked currents were mediated through P2X receptors, we applied trinitrophenyl-ATP (TNP-ATP), a selective homomeric P2X1, homeric P2X3, and heteromeric P2X2/3 receptor antagonist (Virginio et al., 1998). In the presence of TNP-ATP, the αβ-metATP-mediated slowly desensitizing current was dose-dependently antagonized (Fig. 4A, C); the IC50 value of the antagonist was 20.1 ± 6 nM (n = 5). A317941, a potent and selective homomeric P2X3 and heteromeric P2X2/3 receptor antagonist (Jarvis et al., 2002), also attenuated the current (Fig. 4B) in a dose-dependent manner (IC50: 51.3 ± 18.8 nM; n = 3) (Fig. 4C). However, in contrast to TNP-ATP, which at the greatest concentration blocked all of the αβ-metATP-mediated current (1 μM TNP-ATP; 98.4 ± 1%), A317941 only attenuated 34.9 ± 10% of the control αβ-metATP-evoked current at the greatest concentration tested (10 μM). To confirm that the slowly desensitizing
current produced by αβ-metATP was mediated by heteromeric P2X2/3 receptors, we also applied TNP-ATP (100 μM) to every cell that was tested with A317491. Consistent with the description above, TNP-ATP antagonized almost all of the αβ-metATP-evoked current, whereas A317491 only partially blocked the current (Fig. 4A,B). Both TNP-ATP and A317491 completely and reversibly inhibited the fast desensitizing current in all five neurons tested (Fig. 4D,E). Therefore, the slowly and rapidly desensitizing currents studied here are likely to be mediated through heteromeric P2X2/3 and homomeric P2X3 receptors, respectively.

Characterization of proton-sensitive currents in bladder sensory neurons

ATP is an acid, and although 1 mM ATP did not change the pH of external solution buffered with 10 mM HEPES, lowering the buffering capacity of the external solution to 0.05 mM unmasked the acidic nature of ATP (Fig. 4F). Considering the potential of ATP to briefly acify the extracellular space when released, it is possible that bladder afferents may also respond to protons. Therefore, we challenged with protons bladder sensory neurons that were tested previously for purinergic responsiveness.

LS bladder neurons

Application of pH 5 caused an inward current in 42 of 54 (77.8%) LS bladder neurons tested (Fig. 5). We could differentiate transient and sustained currents, with the majority of cells expressing both types in response to protons. Based on activation and desensitization kinetics, we separated transient currents into rapidly and slowly desensitizing currents. Nine of 42 LS bladder neurons (21.4%) expressed fast currents, all of which displayed two peaks with two distinct activation and desensitization time constants (p < 0.01) (Fig. 5A, Table 3). Accordingly, we termed the first and second rapidly desensitizing currents fast and intermediate, respectively, to distinguish them from the slowly desensitizing current. An additional 13 of 42 (31%) proton-sensitive LS bladder neurons showed predominantly slowly activating and slowly desensitizing currents (Fig. 5B). In the remaining 20 of 42 (47.6%) neurons, protons triggered mixed transient and sustained currents (Fig. 5C). The desensitization kinetics of various proton-induced currents is summarized in Table 3. Current densities for the fast, intermediate, and slowly desensitizing and sustained components were 56.4 ± 10.57.8 ± 11.52.2 ± 15.5, and 86.1 ± 16 pA/pF, respectively.

Figure 4. Antagonism of αβ-metATP-evoked currents. A, The αβ-metATP-evoked slowly desensitizing current was significantly attenuated in the presence of TNP-ATP. The inhibition was reversible with complete recovery after washout. B, In the same neuron, A317491 failed to completely antagonize the αβ-metATP-evoked current. The partial inhibition was completely reversible. C, TNP-ATP completely abolished the αβ-metATP-evoked current with an IC50 value of 20.1 ± 6 nM (n = 5), whereas A317491 only partially blocked the αβ-metATP-evoked current (IC50, 51.3 ± 18.8 nM; n = 3). D, Proportion of maximal current. Similarly, the αβ-metATP-evoked fast desensitizing current (left trace) was completely inhibited by TNP-ATP (middle trace) and recovered after washout (right trace). E, In the same cells (n = 5), A317491 at 100 nM (left trace) and 10 μM (middle trace) completely antagonized the αβ-metATP-evoked current with full recovery after washout (right trace). Note that the sustained component was unaffected by A317491.

Figure 5. Proton-evoked currents in bladder sensory neurons. Protons (pH 5) produced a spectrum of currents with distinct desensitizing kinetics: fast, intermediate, and slowly desensitizing and sustained. Three current patterns were identified in LS neurons: fast and intermediate desensitizing (A), slowly desensitizing (B), and fast and intermediate desensitizing (C), followed by sustained. Similarly, fast and intermediate desensitizing (D), slowly desensitizing (E), fast and intermediate desensitizing (F) followed by a sustained current, was found in TL neurons. Protons produced progressively greater currents in a concentration-dependent manner as shown in G, H, I. pH response relationships for various proton-evoked currents in LS (H) and TL (I) neurons. Fast (○), intermediate (□), and sustained (●) currents were pH dependent with EC50 values of 5.79 ± 0.27 (n = 12), 5.48 ± 0.45 (n = 12), and 4.51 ± 0.1 (n = 8), respectively, for LS neurons. Similarly, the EC50 values in TL neurons for fast, intermediate, and sustained currents were 5.89 ± 0.20 (n = 7), 6.19 ± 0.24 (n = 7), and 4.73 ± 0.2 (n = 3), respectively. The EC50 values of the sustained component of the current were significantly lower than those for fast and intermediate currents (p < 0.05). lmax, Proportion of maximal current; I, ATP concentration–pH relationship. In 10 mM HEPES-buffered solution (filled circle; n = 3), ATP up to 1 mM did not change the pH, whereas ATP produced a concentration-dependent drop in pH in 0.05 mM HEPES-buffered solution (open symbol; n = 3).
**Table 3. Summary of proton-evoked current kinetics in bladder sensory neurons**

<table>
<thead>
<tr>
<th></th>
<th>pH 5</th>
<th>tau (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>318 ± 20</td>
<td>311 ± 70</td>
</tr>
<tr>
<td>Intermediate</td>
<td>1056 ± 111</td>
<td>592 ± 84</td>
</tr>
<tr>
<td>Slow</td>
<td>1784 ± 183</td>
<td>1433 ± 331</td>
</tr>
<tr>
<td>Sustained</td>
<td>3935 ± 66</td>
<td>n/a</td>
</tr>
<tr>
<td><strong>TL</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast</td>
<td>370 ± 28</td>
<td>266 ± 37</td>
</tr>
<tr>
<td>Intermediate</td>
<td>1326 ± 98</td>
<td>652 ± 46</td>
</tr>
<tr>
<td>Slow</td>
<td>1603 ± 104</td>
<td>4309 ± 778</td>
</tr>
<tr>
<td>Sustained</td>
<td>3892 ± 88</td>
<td>n/a</td>
</tr>
</tbody>
</table>

tp, Time to peak; n/a, not applicable.

**Figure 6.** Recovery kinetics of proton-evoked fast and intermediate desensitizing currents in LS (A) and TL (B) neurons, respectively. Both currents recovered partially from desensitization after 3 s (left pair of traces) and almost completely after 5 s (right pair of traces). The recovery kinetics of both current types followed a single-exponential function in LS (C) and TL (D) neurons with time constants (fast and intermediate, respectively) of 0.65 ± 0.18 s (n = 4) and 1.1 ± 0.21 s (n = 4) for LS neurons and 0.56 ± 0.13 s (n = 3) and 1.26 ± 0.37 s (n = 3) for TL counterparts.

**TL bladder neurons**

A similar fraction of TL bladder neurons (45 of 52; 86.5%) responded to protons. Based on kinetics, we differentiated transient and sustained currents (Fig. 5D–F). Five (11.2%) neurons expressed transient currents with fast desensitization (Fig. 5D), and an additional 20 (44.4%) neurons responded with only slowly desensitizing currents (Fig. 5E). The remaining 20 cells (44.4%) expressed mixed transient and sustained currents (Fig. 5F). Although no differences in the fraction of neurons with distinct kinetics and kinetics of fast and intermediate desensitizing currents were found in LS neurons compared with their TL counterparts, the slow current in TL neurons desensitized significantly slower than in LS neurons (p < 0.001). Current densities were 48.2 ± 10, 59 ± 12, 74.8 ± 15, and 30.3 ± 7 pA/pF for fast, intermediate, and slowly desensitizing and sustained components, respectively. Whereas the current densities of fast, intermediate, and slowly desensitizing components did not differ, the sustained current density was significantly less in TL than in LS bladder neurons (p < 0.001).

**pH dependency**

Recent evidence suggests that homomeric ASIC1a and ASIC3 channels are most sensitive to protons (Benson et al., 2002; Hes-selager et al., 2004). These homomeric channels are typically characterized by rapid desensitization kinetics similar to the fast and intermediate desensitizing currents described above. To further differentiate proton-evoked currents, we explored proton sensitivity by investigating the pH response relationship in neurons that predominantly expressed fast and intermediate desensitizing currents (Fig. 5G). Lowering the pH from 7 to 4 evoked increasingly larger currents in LS bladder neurons with pH50 values of 5.79 ± 0.27 (n = 12) and 5.48 ± 0.45 (n = 12) for fast and intermediate desensitizing currents, respectively (Fig. 5H). Similarly, progressively larger currents were evoked by increasing the concentration of protons in TL neurons, with pH50 values of 5.89 ± 0.20 (n = 7) and 6.19 ± 0.24 (n = 7) for fast and intermediate desensitizing currents, respectively (Fig. 5I). pH sensitivity did not differ for fast and intermediate desensitizing proton-gated currents in either LS or TL neurons. The sustained proton-activated current in both LS and TL neurons, however, had a low pH threshold that was apparent as the pH was decreased to ≤5.5; pH50 values for LS and TL bladder neurons were 4.51 ± 0.1 (n = 8) and 4.73 ± 0.2 (n = 3), respectively. Although the pH50 values of the sustained component did not differ between LS and TL neurons, they were significantly lower than the pH50 values determined for the fast and intermediate desensitizing currents described above (p < 0.05).

**Recovery kinetics**

We next examined the recovery kinetics of transient proton-evoked currents from desensitization. As described above, we varied the interapplication duration and measured the fractions of current recovered over time. As shown in Figure 6, both fast and intermediate desensitizing currents in LS and TL neurons recovered rapidly and almost completely from desensitization within 5 s. The kinetics of recovery followed a single-exponential function with time constants of 0.65 ± 0.18 s (n = 4) and 1.1 ± 0.21 s (n = 4) for fast and intermediate desensitizing currents, respectively, in LS neurons (Fig. 6C). Similarly, the recovery time constants for fast and intermediate desensitization currents were 0.56 ± 0.13 s (n = 3) and 1.26 ± 0.37 s (n = 3) in TL neurons (Fig. 6D). The recovery kinetics for fast and intermediate desensitizing currents did not differ and were similar for LS and TL neurons.

**Antagonism of proton-evoked currents by amiloride and capsaicin**

To determine whether proton-evoked currents were mediated through ASICs, we applied amiloride (200 μM), a nonselective ASIC blocker (McCleskey and Gold, 1999). In the presence of amiloride, proton-evoked fast and intermediate desensitizing currents were reduced to 1.7 ± 0.7% (n = 5) and 9.8 ± 3.2% (n = 5) of control, respectively, indicating that transient proton-evoked currents were mediated by ASICs. Interestingly, the sustained component of the current was also inhibited to 26.4 ± 8%
TRPV1 channels in the same neurons. Application of capsazepine sustained component of the current was mediated through used. Therefore, we applied the TRPV1 receptor antagonist cap-
account for the sustained component that we saw when pH 5 was at high concentrations can activate TRPV1 channels and could after washout (Fig. 7
Inhibition was reversible after washout. B. In the same neurons, capsazepine inhibited the sustained current evoked by pH 5 to 31 ± 13.3% of control without affecting the fast and intermediate desensitizing currents. Only partial recovery from capsazepine-produced inhibition was observed. All traces were taken from the same neuron. (n = 5) of control. All three components recovered completely after washout (Fig. 7A). Previous studies have shown that protons at high concentrations can activate TRPV1 channels and could account for the sustained component that we saw when pH 5 was used. Therefore, we applied the TRPV1 receptor antagonist cap-sazepine (Dickenson and Dray, 1991) to determine whether the sustained component of the current was mediated through TRPV1 channels in the same neurons. Application of capsazepine (100 μM) reduced the sustained current to 31 ± 13.5% of control without affecting the transient currents, suggesting that the sus-
tained component is mediated, in part, through TRPV1 channels (Fig. 7B).
Effects of capsaicin
Because protons can activate TRPV1 channels, capsaicin was used to further confirm the presence of functional TRPV1 recep-
tors on bladder sensory neurons that were tested previously with purinergic agonists and protons. Capsaicin (1 μM) evoked a per-
sistent inward current in 49 of 54 (90.7%) LS neurons with a current density of 129.6 ± 17 pA/pF (Fig. 8D). All proton-
sensitive neurons (42 of 42) that displayed a transient compo-
nent(s) were also responsive to capsaicin, suggesting colocaliza-
tion of functional ASIC and TRPV1 receptors in LS bladder sensory neurons. In addition, all proton-sensitive neurons with a sustained current (48%; 22 of 42) responded to capsaicin. Seventy-six percent (41 of 54) of LS neurons responded to α,β-
metATP as well as protons and capsaicin, suggesting a high de-
gree of colocalization of functional P2X, ASIC, and TRPV1 recep-
tors on these neurons (Fig. 8). The percentage of bladder sensory neurons that responded to α,β-metATP, protons, and capsaicin is summarized in Table 4.
Similarly, capsaicin produced an inward current in 48 of 52 (92.3%) TL neurons with a current density of 86.9 ± 11 pA/pF (Fig. 8H). Although the percentage of capsaicin-sensitive neu-ons did not differ between LS and TL bladder sensory neurons, the capsaicin-evoked current density was significantly less in TL neurons than in LS counterparts (p < 0.05), consistent with the proton-evoked sustained current being mediated through TRPV1 channels. Similar to LS neurons, all proton-sensitive TL neurons exhibiting a sustained component (44%; 20 of 45) also responded to capsaicin. Almost all proton-sensitive TL neurons (44 of 45) that had a transient component(s) also responded to capsaicin, suggesting a lesser degree of colocalization of P2X, ASIC, and TRPV1 channels in TL than in LS bladder neurons (p < 0.05). We verified immunohistochemically the coexpression of P2X and TRPV1 receptors on bladder sensory neurons that were retrogradely labeled from the bladder by Fluorogold (Fig. 9A,B) but did not attempt to quantify the percentages of receptor expression or coexpression. Consistent with the functional evidence for colocalization of P2X receptors and TRPV1 channels, P2X3 and P2X2 subunits (Fig. 9A) or P2X5 and TRPV1 channels (Fig. 9B) were coimmunostained within the same bladder neurons.
IB4 labeling
Because binding of IB4 is considered to indicate the absence of myelination (Chen et al., 2003), we estimated the number of neurons that give rise to unmyelinated axons in LS and TL DRGs. One coverslip from each rat (n = 3) was selected randomly for cell counting. In control experiments, the binding of IB4 was considered specific because preincubation with unconjugated IB4 (0.2 mg/ml) blocked all binding of the conjugated form (data not shown). Of 2267 LS and 3613 TL neurons, 1071 and 1867, respectively, bound IB4. Thus, IB4-positive neurons comprised 48.8 ± 2.5% (n = 3) and 54.3 ± 3.8% (n = 3) of all LS and TL DRG neurons in the samples studied, respectively (p > 0.05). Approximately 50–70 and 96% of rat bladder pelvic and hypo-
gastric/splanchnic afferents, respectively, have slow conduction velocities consistent with the absence of myelination (Nadelhaft and Vera, 1991; Sengupta and Gebhart, 1994; Shea et al., 2000). To characterize bladder sensory neurons further, we applied IB4 after examination of neurons as described above (Fig. 9C). Sixty-one percent (33 of 54) of LS bladder neurons and 84.6% (44 of 52) of TL bladder neurons were positive for IB4 (χ² = 6.2; p < 0.05). Sixty percent (30 of 50) of α,β-metATP-sensitive LS neu-ons and 84.6% (22 of 26) of α,β-metATP-sensitive TL neurons stained for IB4 (χ² = 0.553; p = 0.4), revealing that not all P2X receptor-expressing cells were IB4 positive. Similarly, most but not all neurons responding to protons were IB4 positive; 31 of 42 (73.8%) and 38 of 45 (84.4%) proton-sensitive LS and TL neu-ons, respectively, were IB4 positive (χ² = 0.9; p = 0.3). Finally,
31 of 49 (63.3%) and 40 of 48 (83.3%) capsaicin-sensitive LS and TL neurons, respectively, stained with IB4 ($\chi^2 = 4.0; p < 0.05$).

**Discussion**

The pelvic and hypogastric/lumbar splanchnic innervations of the bladder subserve different functions (see Introduction). Recent reports have implicated P2X and TRPV1 receptors in micturition (Cockayne et al., 2000; Birder et al., 2002), and we compared in the present study responses of TL and LS bladder sensory neurons to purinergic agonists, protons, and capsaicin. In addition to similarities, we noted differences between the two bladder neuron populations: (1) nearly all LS but only 50% of TL neurons responded to $\alpha,\beta$-metATP; (2) LS and TL neurons exhibited slowly and mixed desensitizing currents, respectively, with LS neurons expressing greater slowly desensitizing current density than TL counterparts; and (3) most bladder neurons also responded to protons and capsaicin, but significantly more LS than TL neurons responded to $\alpha,\beta$-metATP, protons, and capsaicin. We also noted differences in size between TL and LS neurons and between the density of innervation by the two pathways, all of which support differential transmission of sensory information in bladder pelvic and hypogastric/lumbar splanchnicafferent pathways.

Despite the importance of the extrinsic innervation in sensation and micturition (Andersson, 2002), virtually no information is available about the relative densities of pelvic and hypogastric/lumbar splanchnic innervation of the bladder. Acknowledging that we may have underestimated bladder innervation because the entire bladder was not injected with DiI, we found twice as many neurons projecting through the pelvic nerves than the hypogastric/lumbar splanchnic nerves.

Most LS bladder neurons were small or medium in size based on capacitance, a finding consistent with an immunohistochemical study of S1 bladder neurons (Yoshimura et al., 2003) and electrophysiological evidence that 50–70% of pelvic nerve bladder afferents conduct in the C-fiber range (Sengupta and Gebhart, 1994; Shea et al., 2000). In contrast, we found that TL bladder neurons were medium to large in size, although >90% of hypogastric/lumbar splanchnic nerve fibers are unmyelinated based on conduction velocity (Nadelhaft and Vera, 1991). In addition, although P2X3 and TRPV1 are considered to be expressed predominantly on small DRG neurons associated with nociceptors (Caterina et al., 1997, 2000; Vulchanova et al., 1998), the present results question the use of cell size as a surrogate index of axon myelination and reveal that many medium or large bladder sensory neurons also express functional P2X3 and TRPV1 receptors, emphasizing that, at least in visceral sensory neurons, cell diameter is an unreliable indicator of presumed function.

Because IB4 binding is considered a marker for neurofilament-negative neurons associated with unmyelinated axons (Yoshimura et al., 2003), we stained recorded cells for IB4 (Chen et al., 2003). Approximately 50–70 and 96% of pelvic and hypogastric/lumbar splanchnic nerve bladder afferents, respectively, conduct in the C-fiber range (Nadelhaft and Vera, 1991; Sengupta and Gebhart, 1994; Shea et al., 2000), consistent with the absence of neurofilament staining in ~70% of bladder LS neurons (Yoshimura et al., 2003). In support, we found that most TL (85%) and LS (61%) neurons bound IB4. Although these findings are consistent with physiological data, Yoshimura et al. (2003) reported that only 20% of LS bladder neurons bound IB4. The discrepancy may in part be attributable to technical differences; we noted significantly less staining of bladder neurons by FITC–IB4 compared with AlexaFluor 488–IB4 (at the same concentration). Despite the apparent higher sensitivity of AlexaFluor 488–IB4, only ~60% of $\alpha,\beta$-metATP-responsive LS bladder neurons bound IB4 in contrast to the >90% of unlabeled, small-sized P2X3-immunoreactive DRG neurons reported to bind IB4 (Vulchanova et al., 1998). In further contrast, although nearly 90% of TL bladder neurons bound IB4, Su et al. (1986) reported that most TL innervating the urinary tract were calcitonin gene-related peptide immunoreactive. Thus, as appears true for P2X and TRPV1, IB4 may not reliably identify nonpeptidergic bladder neurons. These data highlight differences between bladder (visceral) and somatic sensory neurons, further supporting that visceral and somatic sensory neurons are functionally distinct (McMahon, 1997).

The homomeric P2X3, homomeric P2X3, and heteromeric P2X2/3 receptor agonist $\alpha,\beta$-metATP triggered a slowly desensitizing current in most LS neurons (86%), consistent with a recent report (Zhong et al., 2003). Rapidly inactivating and mixed ki-

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**Figure 8.** Sensitivity to purinergic agonists, protons, and capsaicin in bladder sensory neurons. Sequential application of ATP (A), $\alpha,\beta$-metATP (B), pH 5 (C), and capsaicin (D) to the same neuron evoked inward currents in 75.9% (41 of 52) of LS neurons. Similarly, ATP (E), $\alpha,\beta$-metATP (F), pH 5 (G), and capsaicin (H) produced inward currents in 48.1% (25 of 52) of TL neurons.

**Table 4. Summary of bladder sensory neurons that respond to $\alpha,\beta$-metATP, protons, and capsaicin**

<table>
<thead>
<tr>
<th></th>
<th>$\alpha,\beta$-metATP</th>
<th>H+</th>
<th>Capsaicin</th>
<th>$\alpha,\beta$-metATP and H+</th>
<th>$\alpha,\beta$-metATP and capsacin</th>
<th>H+ and capsacin</th>
<th>$\alpha,\beta$-metATP, H+, and capsacin</th>
</tr>
</thead>
<tbody>
<tr>
<td>LS</td>
<td>92.6% (50/54)</td>
<td>77.8% (42/54)</td>
<td>90.7% (49/54)</td>
<td>84% (42/50)</td>
<td>98% (49/50)</td>
<td>100% (42/42)</td>
<td>75.9% (41/54)</td>
</tr>
<tr>
<td>TL</td>
<td>50% (26/52)*</td>
<td>86.5% (45/52)</td>
<td>92.3% (48/52)</td>
<td>100% (26/26)</td>
<td>100% (26/26)</td>
<td>97.8% (44/45)</td>
<td>48.1% (25/52)*</td>
</tr>
</tbody>
</table>

Data are a comparison between LS and TL neurons ($^*p < 0.05$).
nomic currents were produced by $\alpha,\beta$-metATP in <15% of LS neurons. In contrast, only one-half of TL bladder neurons studied expressed functional P2X receptors, revealing that P2X receptors are differentially expressed in the two afferent bladder pathways. Furthermore, $\alpha,\beta$-metATP triggered rapidly inactivating or mixed currents in most of these TL neurons. The desensitization kinetics of mixed currents followed a double-exponential function, suggesting heterogeneous P2X receptor expression in the same neuron. Based on current kinetics and antagonism by TNP-ATP [a P2X1, P2X3, and P2X2/3 receptor antagonist (Virginio et al., 1998)] and previous reports demonstrating (1) dependence on P2X$_3$ for $\alpha,\beta$-metATP sensitivity (Cockayne et al., 2000), (2) high incidence of colocalization of P2X$_2$ and P2X$_3$ (Vulchanova et al., 1997), and (3) absence of P2X1 receptor transcripts in DRG neurons (Kobayashi et al., 2005), the present data suggest that activation of P2X$_{2/3}$ receptors by $\alpha,\beta$-metATP produces the slowly desensitizing current. Additionally, both TNP-ATP and the P2X$_3$ and P2X$_2/3$ receptor antagonist A317491 (Jarvis et al., 2002) completely antagonized the fast desensitizing current, suggesting that activation of homomeric P2X$_3$ receptors mediates the fast desensitizing current. However, even at high concentrations (10 $\mu$M), A317491 only antagonized ~30% of the slowly desensitizing current. This contrasts with a complete block of the fast desensitizing current at significantly lower concentrations (100 nM), consistent with previous data for the fast current (Jarvis et al., 2002). These findings in native bladder sensory neurons demonstrate greater inhibitory effects of A317491 on homomeric P2X$_3$ compared with heteromeric P2X$_{2/3}$ receptors, consistent with data from heterologous expression systems (Jarvis et al., 2002), and clearly show a more striking difference of inhibitory effects on the kinetically distinct currents. Based on kinetic and pharmacological characteristics, LS bladder neurons express predominantly P2X$_{2/3}$ receptors, whereas heterogeneous populations of homomeric P2X$_3$ and heteromeric P2X$_{2/3}$ receptors are found in TL counterparts. Because (1) significantly more LS than TL neurons responded to purinergic agonists, (2) LS neurons expressed significantly greater peak current density (slow type) than TL counterparts, and (3) the differences in desensitization and recovery kinetics, we suggest that purinergic transmission under normal physiological conditions is predominantly conveyed through pelvic rather than the hypogastric/lumbar splanchnic afferents by activating heteromeric P2X$_{2/3}$ receptors.

Because ATP is acidic, ATP released in sufficient quantities could acidify the immediate microenvironment and activate P2X as well as proton-sensitive channels. In support, most (>80%) $\alpha,\beta$-metATP-sensitive neurons also responded to protons, suggesting a high degree of colocalization of ASIC and P2X channels. ASIC channels are implicated because proton sensitivity of the cells is relatively high and amiloride, an ASIC channel antagonist (McCleskey and Gold, 1999), antagonized effects. Based on the presence of multiple components and complex current kinetics, our data suggest the existence of multiple ASIC subunits in the same cell. The transient currents had similar pH$_{50}$ values, consistent with previous findings showing that most ASIC channels in vivo are heteromers that show no difference in proton sensitivity compared with homomeric forms (Benson et al., 2002; Hesselager et al., 2004). In contrast, the sustained component of the proton-gated current showed significantly lower pH$_{50}$ compared with pH$_{50}$ values determined for transient components and was partially antagonized by amiloride and capsazepine, a TRPV1 receptor antagonist (Dickenson and Dray, 1991). Although transient and sustained proton-activated currents were indiscriminately antagonized by amiloride, capsazepine selectively suppressed the sustained component. Based on lower proton sensitivity, current kinetics, and antagonism by capsazepine, activation of TRPV1 receptors, in part, accounts for the sustained current in bladder neurons.

Because TRPV1 receptors are important to bladder function (Caterina et al., 1997, 2000; Tominaga et al., 1998; Birder et al., 2002), we examined bladder neuron sensitivity to capsaicin. More than 90% of bladder neurons responded to capsaicin with a sustained current, confirming the presence of functional TRPV1 receptors, which have been documented immunohistochemically (Cockayne et al., 2000; Birder et al., 2002). One might expect bladder neurons that responded to capsaicin to also exhibit a sustained current to protons, but neither we nor others (Stein et al., 1992; Dirajlal et al., 2003) found such a relationship. Both the sustained proton-evoked current and capsaicin-activated peak current densities were greater in LS than in TL neurons. Moreover, significantly more LS neurons (76%) responded to $\alpha,\beta$-metATP, protons, and capsaicin than did TL counterparts (48%), suggesting a greater degree of P2X, ASIC, and TRPV1 receptor colocalization in

**Figure 9.** Immunohistochemistry of bladder sensory neurons. A, Triple immunohistochemical staining of P2X receptors in bladder sensory neurons identified by the presence of Fluorogold (FG), P2X$_2$, and P2X$_3$. The merged image demonstrates the colocalization of P2X$_2$ and P2X$_3$ subunits in a bladder sensory neuron. B, Similarly, a bladder sensory neuron was identified by the presence of FG, P2X$_2$, and TRPV1. The merged image shows coexistence of P2X$_2$ and TRPV1 receptors in a bladder sensory neuron. C, IB4 labeling in bladder sensory neurons. A bladder sensory neuron was identified by the presence of Dil (left) and bound to IB4 (middle). The merged image confirms the bladder sensory neuron was positive for IB4 (right).
LS than in TL bladder neurons. Consistent with these functional data, coexpression of the receptor proteins on bladder sensory neurons was also seen immunohistochemically.

**Significance of functional P2X, ASIC, and TRPV1 channel colocalization**

Release of ATP from the urothelium during bladder distension contributes to mictrition (Cockayne et al., 2000; Vlaskovska et al., 2001), and Burnstock (1996) has suggested that ATP within hollow organs can mediate painful sensations. In support of the latter, significantly more ATP is released from urothelial cells in interstitial cystitis patients compared with normal subjects (Sun et al., 2001). The high degree of colocalization of functional P2X, ASIC, and TRPV1 channels in bladder neurons reported here suggests potential interaction/modulation and cotransmission by ATP and protons. ATP is acidic, and the potential release of three additional protons through the hydrolysis of ATP to adenosine could further acidify the microenvironment (Robergs et al., 2004). Of the seven known P2X receptor subunits, heteromeric P2X<sub>2/3</sub>- or homomeric P2X<sub>2</sub>-mediated currents are potentiated by protons, whereas the homomeric P2X<sub>2</sub>-mediated current is inhibited by acidic pH (Stoop et al., 1997). Accordingly, we propose that ATP, by virtue of its acidic nature, could also activate ASIC and TRPV1 channels. The net effect would result in larger currents associated with greater depolarization and an increase in action potential number that may partly account for altered sensory transmission and sensitization of the bladder after cystitis.

**References**


