Spontaneous Electrical Activity of Cultured Interstitial Cells of Cajal from Mouse Urinary Bladder

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Interstitial cells of Cajal (ICCs) from the urinary bladder regulate detrusor smooth muscle activities. We cultured ICCs from the urinary bladder of mice and performed patch clamp and intracellular Ca\(^{2+}\) ([Ca\(^{2+}\)]\text{c}) imaging to investigate whether cultured ICCs can be a valuable tool for cellular functional studies. The cultured ICCs displayed two types of spontaneous electrical activities which are similar to those recorded in intact bladder tissues. Spontaneous electrical activities of cultured ICCs were nifedipine-sensitive. Carbachol and ATP, both excitatory neurotransmitters in the urinary bladder, depolarized the membrane and increased the frequency of spike potentials. Carbachol increased [Ca\(^{2+}\)]\text{c} oscillations and basal Ca\(^{2+}\) levels, which were blocked by atropine. These results suggest that cultured ICCs from the urinary bladder retain rhythmic phenotypes similar to the spontaneous electrical activities recorded from the intact urinary bladder. Therefore, we suggest that cultured ICCs from the urinary bladder may be useful for cellular and molecular studies of ICCs.

Key Words: Interstitial cells of Cajal, Spontaneous electrical activities, Urinary bladder

INTRODUCTION

Interstitial cells of Cajal (ICCs) play an important role regulating smooth muscle cells by generating spontaneous pacemaker potentials and mediating signals from enteric neurons to smooth muscle cells as well as serving as stretch sensors in the gastrointestinal tract [1,2]. ICCs are distributed throughout the urinary tract and play an important role regulating smooth muscle activities [3,4]. Several types of ICCs have been found throughout the urinary bladder in several species [5,6]. ICCs are c-kit positive cells and spontaneous phasic contractions of the urinary bladder are inhibited by c-kit blocker [7]. The urinary bladder is innervated with cholinergic, peptidergic, and nitric oxide nerve fibers, revealing close relationship with ICCs, and indicating that urinary bladder ICCs regulate smooth muscle activities and mediate neural signals to smooth muscles [6,8]. It has been reported that enhanced spontaneous phasic contractions in overactive bladder may be caused by release of acetylcholine (ACh) from cholinergic nerves acting on muscarinic receptors in ICCs [9]. ICCs have mechanical sensitivity; thus, they contribute to mechanosensitive conductance during urinary bladder regulation [10], and the distribution of ICCs is altered in urinary bladder outlet obstruction (BOD) [11]. These results suggest that ICCs play an important role in urinary bladder physiology and pathology. Therefore, ICC investigations might benefit the understanding of urinary bladder pathophysiology. Although the physiological roles of ICCs have been demonstrated, many studies have been morphological using immunohistochemistry and electrophysiology of intact tissues. However, because intact urinary bladder tissues contain other cells (smooth muscle cells and neuronal cells), it is difficult to identify the pure functional and physiological roles of ICCs alone. Cultured ICCs from the gastrointestinal tract have spontaneous pacemaker potentials similar to slow waves recorded in intact tissues; therefore, cultured ICCs have been used to elucidate the functional role of ICCs [12-14]. In the present study, we investigated whether cultured ICCs from urinary bladder can be of benefit to understand the functional physiological roles like gastrointestinal ICCs. Thus, we cultured urinary bladder ICCs from mice and recorded spontaneous electrical potentials.

ABBREVIATIONS: ICCs, interstitial cells of Cajal; BOD, bladder outlet obstruction.
METHODS

Preparation of cells

All experiments were performed according to the Guiding Principles for the Care and Use of Animals approved by the Ethics Committee of Chosun University and the National Institutes of Health Guide for the Care and Use of Laboratory Animals. Every effort was made to minimize both the number of animals used and their suffering. Balb/C mice (8–13 days old) of either sex were anesthetized with ether and sacrificed by cervical dislocation. The urinary bladder was removed and opened. The luminal contents were washed away with Krebs-Ringer bicarbonate solution. The tissues were pinned to the base of a Sylgard dish and the mucosa was removed by sharp dissection. Small strips of bladder tissues were equilibrated in Ca$^{2+}$-free Hank’s solution containing 5.36 mM KCl, 125 mM NaCl, 0.34 mM NaOH, 0.44 mM Na$_2$HCO$_3$, 10 mM glucose, 2.9 mM sucrose, 11 mM HEPES for 30 min, and the cells were dispersed with an enzyme solution containing 1.3 mg/ml collagenase (Worthington Biochemical, Lakewood, NJ USA), 2 mg/ml bovine serum albumin (Sigma, St. Louis, MO USA), 2 mg/ml trypsin inhibitor (Sigma) and 0.27 mg/ml ATP. Cells were plated onto sterile glass coverslips coated with murine collagen (2.5 μg/ml, Falcon/BD, Franklin Lakes, NJ USA) in 35 mm culture dishes. The cells were then cultured at 37°C in a 5% CO$_2$ incubator in smooth muscle growth medium (Clonetics, San Diego, CA USA) supplemented with 2% antibiotic/antimycotic (Gibco, Grand Island, NY USA) and murine stem cell factor (5 ng/ml; Sigma). ICCs were identified immunologically with the use of a monoclonal antibody for kit protein (ACK2) labeled with Alexa Fluor 488 (Molecular Probes, Eugene, OR, USA).

Labeling of cultured ICCs by c-kit immunofluorescence

Cultured ICCs was fixed in acetone (20°C/5 min), and the preparations were washed for 60 min in phosphate-buffered saline (PBS; 0.01 M, pH 7.4). Cultured ICCs were then incubated in 10% goat serum containing 1% bovine serum albumin for 1 h at room temperature to reduce nonspecific antibody binding. To examine the ICCs, cultured ICC were incubated overnight at 4°C with a rat monoclonal antibody raised against Kit protein (ACK2; 5 μg/ml; Gibco-BRL, Gaithersburg, MD, U.S.A). Immunoreactivity was detected using fluorescein isothiocyanate (FITC)-conjugated secondary antibody (Vector Laboratories Burlingame, CA, USA), 1 × 100 in PBS, 1 h, at room temperature). Control cultured ICCs were prepared in a similar manner, but omitting ACK2 from the incubation medium. The cells were examined under a confocal laser scanning microscope (FV300, Olympus, Tokyo, Japan) at an excitation wavelength appropriate for FITC (488 nm).

Patch clamp experiments

The whole-cell configuration of the patch-clamp technique was used to record membrane potentials (current clamp) from cultured ICCs after 2–3 days of culture. Currents or potentials were amplified by Axopatch 1-D and 200B (Axon Instruments, Foster City, CA, USA). A command pulse was applied using an IBM-compatible personal computer and pClamp software (ver. 6.1; Axon Instruments). The data were filtered at 5 kHz and displayed on an oscilloscope and a computer monitor. Results were analyzed using pClamp and Graph Pad Prism (ver. 2.01) software. All experiments were performed at 30°C.

Solutions and drugs

The cells were bathed in a solution containing 5 mM KCl, 135 mM NaCl, 2 mM CaCl$_2$, 10 mM glucose, 1.2 mM MgCl$_2$, and 10 mM HEPES, adjusted to pH 7.2 with Tris. The pipette solution contained 20 mM K-aspartate, 120 mM KCl, 5 mM MgCl$_2$, 2.7 mM K$_2$ATP, 0.1 mM Na$_2$GTP, 2.5 mM creatine phosphate disodium, 5 mM HEPES, 0.1 mM EGTA, adjusted to pH 7.2 with Tris.

The drugs used were carbachol, methoctramine, glyco- pyrrolate, ATP, and nifedipine. All drugs were purchased from Sigma.

Measurement of intracellular Ca$^{2+}$ concentration

Changes in intracellular Ca$^{2+}$ concentration ([Ca$^{2+}$]) were monitored using fluo-3/AM, which was initially dissolved in dimethyl sulfoxide and stored at −20°C. Cultured ICCs on coverslips (25 mm) were rinsed twice with a bath solution (5 mM KCl, 135 mM NaCl, 2 mM CaCl$_2$, 10 mM glucose, 1.2 mM MgCl$_2$ and 10 mM HEPES, adjusted to pH 7.4 with Tris). The coverslips were then incubated in the bath solution containing 5 μM fluo-3 with 5% CO$_2$ at 37°C for 5 min, rinsed two more times in the bath solution, mounted on a perfusion chamber, and were scanned every 0.4 seconds with a Nikon Eclipse TE200 inverted microscope equipped with a Perkin-Elmer Ultraview confocal scanner and a Hamamatsu Orca ER 12-bit CCD camera (×200). Fluorescence was excited at a wavelength of 488 nm, and emitted light was observed at 515 nm. The temperature of the perfusion chamber containing the cultured ICCs was kept at 30°C during scanning of the Ca$^{2+}$ imaging. Variations in intracellular Ca$^{2+}$ fluorescence emission intensity were expressed as F1/F0, where F0 is the intensity of the first image.

Statistical analysis

Data are expressed as mean±standard error. Differences in the data were evaluated by Student’s t test. p values less than 0.05 were considered as a statistically significant difference. The n values refer to the number of cells used in the patch-clamp experiments.

RESULTS

Spontaneous electrical activities in cultured bladder ICCs

The morphology of cultured ICCs was c-kit positive stellate or spindle-shaped and they were connected to each other (Fig. 1). We performed the current clamp technique with cultured ICCs. The cultured ICCs exhibited two types of spontaneous electrical activities. One was individual generation of spontaneous spike potentials (Fig. 2A). The other was bursting generation of spontaneous spike potentials (Fig. 2B). The resting membrane potential was −41.5±2.7 mV (n=7), the frequency was −2.7±0.5 cycles/5 min (n=7), and the amplitude was 37.7±4.7 mV (n=7) for individual
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Fig. 1. Cultured ICCs from the mouse urinary bladder. The tunica muscularis of the urinary bladder was digested with collagenase, and the dispersed cells were cultured for 2 days. Confocal microscope image of Kit-immunopositive ICCs showed networks in culture.

Fig. 2. Spontaneous electrical activities in cultured ICCs from mouse urinary bladder. The cultured ICCs exhibited two types' spontaneous electrical activities. One is individual generation of spontaneous spike potentials (A). The other is bursting generation of spontaneous spike potentials (B). Dotted lines indicate resting level of membrane potentials.

Fig. 3. Effect of nifedipine on spontaneous electrical activities in cultured ICCs from mouse urinary bladder. In cultured ICCs generated spontaneous spike potentials. Nifedipine (10⁻⁶ M) abolished the spontaneous spike potentials in both individual (A) and bursting (B) types. Dotted lines indicate resting level of membrane potentials.

Fig. 4. Effects of carbachol spontaneous electrical activities in cultured ICCs from mouse urinary bladder. Carbachol (10⁻⁸ M) produced depolarization of the resting membrane with an increase of the frequency of spontaneous spike potentials in individual type (A). In the bursting type, carbachol produced depolarization of the resting membrane with a decrease of the frequency of spontaneous spike potentials (B). Atropine (10⁻⁶ M) blocked the carbachol-induced effects on spontaneous spike potentials in both individual (C) and bursting (D) types. Dotted lines indicate resting level of membrane potentials. CCh: carbachol.

spontaneous spike potentials. The resting membrane potential for bursting spontaneous spike potentials was −42.7±−5.1 mV (n=8), the frequency was mean 7.5±1.5 cycles/5 min (n=8), and the amplitude was 52.5±6.3 mV (n=8). The summarized data are shown in Fig. 2C-E. The application of nifedipine (L-type Ca²⁺ channel blocker; 10⁻⁶ M) almost blocked the spontaneous electrical activities in both types (Fig. 3) (n=5 in individual type and n=4 in bursting type, respectively).

**Effects of carbachol and ATP on spontaneous electrical activities**

Exposure to carbachol (10⁻⁸ M), a selective muscarinic receptor agonist, produced depolarization of the resting membrane with an increase of the frequency of spontaneous electrical activities of the individual type (Fig. 4A). In the presence of carbachol, the resting membrane potential changed from −38.7±2.1 mV to −22.5±3.7 mV and frequencies was changed from 1.8±0.5 cycles/5 min to 6.1±1.3 cycles/5 min in the individual type (n=4). However, carbachol produced depolarization of the resting membrane in the bursting type with a decrease in the frequency of spontaneous electrical activities (Fig. 4B, n=4). In the presence of carbachol, the resting membrane potential changed from −37.3±3.2 mV to −10.7±4.7 mV and the frequencies changed from 9.1±1.1 cycles/5 min to 2.3±1.3 cycles/5 min in the bursting type (n=3). The carbachol-induced effects were blocked by atropine (a muscarinic receptor antagonist; 10⁻⁶ M) in both types (Fig. 4C and 4D), indicating that muscarinic receptors act on ICCs (Fig. 4C). We also investigated ATP effects on spontaneous electrical activities in
two types of spontaneous electrical activities during intractions. In previous studies, detrusor smooth muscle showed ICs contain several ion channels, receptors, and signal activities.

and increased the frequency of spontaneous electrical transient depolarization with mechanical contrac-

tions in the urin ary bladder, depolarized the membrane and increases spontaneous phasic contractions of de-

trusor smooth muscles through ICCs after stimulation by nifedipine, suggesting that L-type Ca2+ channels are crit-

cal to generate the spontaneous electrical activity in u-

rin ary bladder ICCs. L-type Ca2+ channels have been re-

corded in isolated ICCs from the urinary bladder [23]. These results suggest that cultured ICCs retain the sponta-

neous electrical phenotype that similar to that recorded from intact tissues.

The urinary bladder is innervated by the autonomic nerv-

ous system. Parasympathetic innervations induce detrusor smooth muscle contraction with ACh after binding to muscarinic receptors [24]. ATP is also co-released with ACh from parasympathetic nerves and released from urothelial cells during bladder distension; thus, inducing contraction of detrusor muscles after binding to purinergic receptors [25,26]. In contrast, sympathetic innervation and nitric oxide induce relaxation via β-receptors and the cGMP path-

way, respectively [27,28]. Bladder ICCs are innervated with cholinergic nerve fibers and the neurogenic-induced detrusor smooth muscle contractions are blocked by a c-kit blocker, indicating that bladder ICCs mediate neural sig-

nals to detrusor muscles [7]. ACh depolarizes the mem-

brane and increases spontaneous phasic contractions of det-

rusor smooth muscles [21]. In the present study, carbachol, a muscarinic receptor agonist, depolarized the membrane and increased the frequency of spike potentials. Furthermore, the potentials were blocked by atropine, suggesting that cholinergic stimulation modulates detrusor smooth muscle contractions through ICCs after stimulation by muscarinic receptors. The depolarization by carbachol was prominent in spontaneous bursting type spike potentials, whereas increased frequency was prominent in the individual type of spike potentials. We do not know the differ-

Fig. 6. The regulation of carbachol on spontaneous Ca2+ oscillation in cultured ICCs from the mouse urinary bladder. (A) Sequential fluorescence intensity images of Fluo-3 loaded cultured ICCs by carbachol (10^-8 M). In the presence of carbachol, the [Ca(2+)]i oscillations were increased. (B) Sequential fluorescence intensity images of Fluo-3 loaded cultured ICCs by carbachol in the presence of atropine (10^-6 M). Atropine blocked the carbachol-induced effects on [Ca(2+)]i oscillations. CCh, carbachol.

cultured ICs. ATP showed very similar effects as carba-

chol. Exposure to ATP (5×10^-8 M) produced depolarization of the resting membrane with an increase in the frequency of spontaneous electrical activities in the bursting type, indicating that purinergic transmission occurs in bladder ICCs (Fig. 5A). The summarized data are shown in Fig. 5B and 5C.

Effects of carbachol on intracellular Ca2+ intensity in cultured ICs

As intracellular Ca2+ ([Ca(2+)]i) oscillations in ICs are the primary mechanism for pacemaking activity in the gastro-

intestinal tract, we examined the effects of carbachol on [Ca(2+)]i oscillations in ICs. Spontaneous [Ca(2+)]i oscillations were observed in a time series from ICs loaded with fluo3-AM. In the presence of carbachol (10^-8 M), the [Ca(2+)]i oscillations increased and this effect was blocked by atropine (10^-6 M) (Fig. 6, n=3).

DISCUSSION

We showed that the cultured ICs from mouse urinary bladder exhibited spontaneous electrical activities which were similar to that those recorded from intact tissues. In addition, carbachol and ATP, both excitatory motor trans-

mitters in the urinary bladder, depolarized the membrane and increased the frequency of spontaneous electrical activities.

ICCs contain several ion channels, receptors, and signal molecules, indicating that ICs can be a therapeutic target in urinary bladder similar to the gastrointestinal tract [15-18]. It has been reported that ICs populations increase in BOD, indicating that bladder ICs play an important role in voiding [19]. Detrusor smooth muscles display spontaneous transient depolarization with mechanical contrac-

tions. In previous studies, detrusor smooth muscle showed two types of spontaneous electrical activities during intra-

cellular microelectrode recording. Several detrusor smooth muscle cells show individually generated spontaneous single spike-shaped potentials, whereas other cells generate spontaenous bursting of several spike-shaped potentials [20,21]. In this study, cultured ICs were c-kit positive stel-

late or spindle in shapes and formed networks with each others. The cultured ICs also displayed two types of sponta-

eous electrical activity, individually generated sponta-

eous single spike-shaped potentials and bursting potentials which were similar to those recorded in intact urinary bladder tissues, suggesting that ICs may act as pace-

making cells in the urinary bladder. Spontaneous electrical activities in the intact urinary bladder are abolished by L-type Ca2+ channels blockers [22]. We found that the sponta-

eous electrical activities of cultured ICs were blocked by nifedipine, suggesting that L-type Ca2+ channels are crit-

cal to generate the spontaneous electrical activity in u-

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The urinary bladder is innervated by the autonomic nerv-

ous system. Parasympathetic innervations induce detrusor smooth muscle contraction with ACh after binding to muscarinic receptors [24]. ATP is also co-released with ACh from parasympathetic nerves and released from urothelial cells during bladder distension; thus, inducing contraction of detrusor muscles after binding to purinergic receptors [25,26]. In contrast, sympathetic innervation and nitric ox-

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more, the potentials were blocked by atropine, suggesting that cholinergic stimulation modulates detrusor smooth muscle contractions through ICs after stimulation by muscarinic receptors. The depolarization by carbachol was prominent in spontaneous bursting type spike potentials, whereas increased frequency was prominent in the individual type of spike potentials. We do not know the differ-
ences in the effects of carbachol now. Several types of ICCs are distributed in urinary bladder. Therefore we think that this different effect by carbachol may be due to the ICC cell type. However, we should study further to identify the mechanism of muscarinic receptor stimulation. ATP is another excitatory neurotransmitter that mediates atropine-resistant fast excitatory junction potentials via purinergic receptors in the urinary bladder [29]. ATP evokes a transient depolarization in the intact urinary bladder tissues [22]. In the present study, external ATP increased the frequency of spike potentials and depolarized the membrane in cultured ICCs, which was very similar to that reported in intact urinary bladder tissues [29]. This means that ATP can modulates bladder activity through ICCs.

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\text{[Ca}^{2+}]\text{], oscillations are essential to generate pacemaker potentials in small intestinal ICCs. Pacemaking in the gastrointestinal tract is initiated by release of Ca}^{2+} \text{ from the endoplasmic reticulum, and is followed by reuptake of Ca}^{2+} \text{ into mitochondria [12]. Urinary bladder ICCs are capable of generating spontaneous Ca}^{2+} \text{ transients, suggesting that they might be essential for generating spontaneous excitation and exposure to carbachol produced Ca}^{2+} \text{ waves [30]. Carbachol increases firing Ca}^{2+} \text{ transients in dispersed urinary bladder ICCs. In the present study, carbachol increased the Ca}^{2+} \text{ oscillations, and these calcium-induced effects were blocked by atropine, suggesting that carbachol effects on [Ca}^{2+} \text{], are correlated with effects obtained from spontaneous electrical activities by carbachol.}
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In conclusion, our cultured ICCs showed similar spontaneous electrical activities to those recorded from intact urinary bladder preparations and they maintained the spontaneous electrical phenotype. Therefore, we think that use of cultured ICCs may be a beneficial method for cellular and molecular pathophysiological studies of the urinary bladder.

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