PIP₂ Activates TRPV5 and Releases Its Inhibition by Intracellular Mg²⁺

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The transient receptor potential type V5 channel (TRPV5) is a Ca²⁺-selective TRP channel important for epithelial Ca²⁺ transport. Intracellular Mg²⁺ causes a fast voltage-dependent block of the TRPV5 channel by binding to the selectivity filter. Here, we report that intracellular Mg²⁺ binding to the selectivity filter of TRPV5 also causes a slower reversible conformational change leading to channel closure. We further report that PIP₂ activates TRPV5. Activation of TRPV5 by PIP₂ is independent of Mg²⁺. Yet, PIP₂ decreases sensitivity of the channel to the Mg²⁺-induced slow inhibition. Mutation of aspartate-542, a critical Mg²⁺-binding site in the selectivity filter, abolishes Mg²⁺-induced slow inhibition. PIP₂ has no effects on Mg²⁺-induced voltage-dependent block. Thus, PIP₂ prevents the Mg²⁺-induced conformational change without affecting Mg²⁺ binding to the selectivity filter. Hydrolysis of PIP₂ via receptor activation of phospholipase C sensitizes TRPV5 to the Mg²⁺-induced slow inhibition. These results provide a novel mechanism for regulation of TRP channels by phospholipase C-activating hormones via alteration of the sensitivity to intracellular Mg²⁺.

INTRODUCTION

Transient receptor potential (TRP) channels are widespread and play many important functions, ranging from thermal, tactile, taste, osmolar and fluid flow sensing, and embryo development to epithelial Ca²⁺ and Mg²⁺ transport (Hoenderop et al., 2002; Montell et al., 2002; Clapham, 2003). They are classified into TRPC, TRPV, TRPM, TRPP, TRPML, TRPA, and TRPN subfamilies (Hoenderop et al., 2002; Montell et al., 2002; Clapham, 2003). The TRPV subfamily is named after the first mammalian member of the subfamily, vanilloid receptor 1 (Montell et al., 2002; Clapham, 2003). TRPV5 and TRPV6 are highly Ca²⁺-selective TRP channels that mediate trans-epithelial Ca²⁺ transport in kidney and intestine (Hoenderop et al., 2002).

Intracellular Mg²⁺ is a cofactor for many enzymes and controls inward rectification of ion channels by causing voltage-dependent block of outward currents (Romani and Scarpa, 2000; Hille, 2001). Recently, intracellular Mg²⁺ has also been reported to regulate the activity of several TRP channels including TRPM6, TRPM7, and TRPV6 via a mechanism different from voltage-dependent block (Nadler et al., 2001; Runnels et al., 2001; Voets et al., 2003, 2004a). The intracellular free Mg²⁺ concentration in most mammalian cells is between 0.5 and 1 mM and changes only slightly in response to physiological stimuli (Romani and Scarpa, 2000). TRPM7 is permeable to Mg²⁺ (Schmitz et al., 2003). It is believed that Mg²⁺ ions entering cells through TRPM7 feedback on the channel to regulate its activity (Schmitz et al., 2003). TRPV6 does not conduct Mg²⁺ in physiological conditions. The physiological role of intracellular Mg²⁺ regulation of TRPV6 is unknown.

PIP₂ directly regulates inward rectifying K⁺ and other channels (Hilgemann et al., 2001). How membrane lipids alter channel function is an area of intense research interests. For many ion channels, direct interaction with PIP₂ stabilizes channels in certain conformations (Hilgemann et al., 2001). By stabilization in one conformation, PIP₂ modulates the responses of ion channels to regulators, such as GTP binding proteins, intracellular pH, Na⁺, ATP, etc. (Huang et al., 1998; Shyng and Nichols, 1998; Sui et al., 1998; Leung et al., 2000).

In this report, we show that intracellular Mg²⁺ causes a fast voltage-dependent block and a slower inhibition of TRPV5. We further show that PIP₂ activates TRPV5 and decreases sensitivity of the channel to the Mg²⁺-induced slow inhibition. Hydrolysis of PIP₂ via receptor activation of phospholipase C increases the sensitivity of TRPV5 to the Mg²⁺-induced slow inhibition. These results provide a novel mechanism for regulation of TRP channels by phospholipase C-activating hormones by increasing the sensitivity to intracellular Mg²⁺.

MATERIALS AND METHODS

Molecular Biology and Cell Culture

Nucleotide coding sequence of cDNAs for rabbit TRPV5 in pCDNA3 mammalian expression vector (Yeh et al., 2003) was used as template for site-directed mutagenesis using a commer-

Abbreviations used in this paper: DCT, distal convoluted tubule; DVF, divalent-free; TRP, transient receptor potential; WMN, wortmannin.
Electrophysiological Recordings

Cell-attached, inside-out, and whole-cell recordings were performed using an Axopatch 200B patch-clamp amplifier (Axon Instruments) as described previously (Yeh et al., 2003, 2005). For cell-attached recordings, the pipette and bath solution contained (in mM) 140 NaAsp (sodium aspartate), 10 NaCl, 1 EDTA, 10 HEPES (pH 7.4), and 140 KAsp, 10 NaCl, 1 EDTA, 10 HEPES (pH 7.4), respectively. For cell-attached recordings shown in Fig. 8 A, the pipette solution contained (in mM) 140 NaCl, 1 MgCl₂, 1 CaCl₂, and 10 HEPES (pH 7.4). For inside-out recordings, membrane patches were excised into a bath solution containing (in mM) 140 NaAsp, 10 NaCl, 10 HEPES (pH 7.4), 10 EDTA, and various amounts of MgCl₂ titrated to the desired concentrations of ionized Mg²⁺ as specified. Bath solution was perfused at ~200 µl per second. The volume of perfusion chamber and connecting tubing is ~100 µl. Thus, it takes 2–3 s to exchange the bath solution. Concentrations of ionized Mg²⁺ were calculated using a computer program based on algorithms by Fabiato (Fabiato, 1988) (www.stanford.edu/~cpatton/webmaxc/webmaxcE.htm). For whole-cell recordings, the initial pipette and bath solution contained (in mM) 140 NaAsp, 10 NaCl, 10 HEPES (pH 7.4), 10 EDTA, and 140 NaAsp, 10 NaCl, 1 EDTA, 10 HEPES (pH 7.4), respectively. For whole-cell recordings shown in Fig. 9 B, the bath solution contained (in mM) 140 NaCl, 1 MgCl₂, 1 CaCl₂, and 10 HEPES (pH 7.4). Intracellular Mg²⁺ in whole-cell recording was altered by intrapipette perfusion of solutions containing (in mM) 140 NaAsp, 10 NaCl, 10 HEPES (pH 7.4), 10 EDTA, and various amount of MgCl₂ titrated to the desired concentrations of ionized Mg²⁺ as specified. The intrapipette perfusion was performed via a capillary quartz tubing (flamed and pulled manually to outer and inner diameter ~50 and 20 µm, respectively) placed within ~200 µm of the tip of the opening (Hilgemann and Lu, 1998). Voltage protocol for each experiment was described in the individual figure. Currents were low-pass filtered at 1 kHz using an 8-pole Bessel filter, sampled every 0.1 ms (10 kHz) with Digidata-1300 interface and stored directly onto computer hard disk using pCLAMP9 software. Data were transferred to compact discs for long-term storage.

Data Analysis

Dose–response curves for inhibition of TRPV5 by intracellular Mg²⁺ and concentration of Mg²⁺ for half-maximal inhibition (IC₅₀) were obtained by fitting relative currents (normalized to maximal currents; I/Imax) at different Mg²⁺ concentrations using the SigmaPlot program according to equation I/Imax = IC₅₀/IC₅₀ + [Mg²⁺]. The fractional electrical distance of the Mg²⁺ binding site from the outer surface of the membrane (R) was calculated according to equation IC₅₀(V) = IC₅₀(0)exp(28RT/ν) (Nilius et al., 2000), where IC₅₀(V) and IC₅₀(0) represent IC₅₀ at the test voltage V and at 0 mV, respectively, and R, T, and F have their usual meanings. Current traces were fitted to a single or double exponential curve (as indicated) to analyze time constant (τ) for inhibition of currents by Mg²⁺. The on-rate constant (Kₘ) for slow inhibition by Mg²⁺ was calculated according to the equation Kₘ = 1/τ[Mg²⁺]. Data are shown as mean ± SEM of number of observations. Statistical comparison was made using unpaired Student’s t test.

RESULTS

Multiple Effects of Intracellular Mg²⁺ on TRPV5: Slow Reversible Inhibition, Fast Voltage-dependent Block, and “Run-down” via Reduction of PIP₂

Effect One: Slow Reversible Inhibition. We studied TRPV5 channels expressed in Chinese hamster ovary (CHO) cells by cell-attached and inside-out patch-clamp recording (Fig. 1 A). In cell-attached recording, TRPV5-expressing CHO cells exhibited the characteristic strongly inwardly rectifying Na⁺ currents (Nilius et al., 2001; Hoenderop et al., 2003; Yeh et al., 2003) (Fig. 1 B, C/A). Inward (closed circles) and outward currents (open circles) both increased markedly when inside-out patch membranes containing TRPV5 channels were excised (Fig. 1 B, I/O). Reversibly inhibition occurring over tens of seconds by intracellular Mg²⁺. It was expected to distinguish it from the much faster (also reversible) voltage-dependent block described below.

Effect Two: Voltage-dependent Block and Distinction from the “Slow Reversible Inhibition”. Intracellular free Mg²⁺ also causes reversible voltage-dependent block of TRPV5 occurring over a relatively much faster time course (in millisecond time scale; see Hoenderop et al., 2003; Vosets et al., 2003). The distinction between the above slow reversible inhibition occurring over tens of seconds and the voltage-dependent block by Mg²⁺ is rather evident from our results (see ramp I-V curves in Fig. 1 C). First, as mentioned above, inward currents shown in Fig. 1 B were measured at the end of 40-ms hyperpolarization (~100 mV) step (indicated by black arrow in voltage ramp protocol in Fig. 1 A). At these time points, Mg²⁺ ions were completely dissociated (from channels) by the hyperpolarization (Fig. 1 C) and thus did not exert voltage-dependent block on channels. Second, the kinetics of voltage-dependent block distinguishes itself from the slow reversible inhibition. The Mg²⁺-induced voltage-dependent block of TRPV5 was evident by the characteristic unblock during the ~100 mV hyperpolar-
multiple effects of intracellular Mg$^{2+}$ on TRPV5. (A) Configuration and voltage protocol of recording. Voltage stimuli were applied every 10 s. Black and gray arrows indicate time points when inward and outward currents shown in Fig. 1 B were taken, respectively. (B) Intracellular Mg$^{2+}$ causes a fast reversible voltage-dependent block, a slow reversible inhibition, and an irreversible run-down of TRPV5 in inside-out membranes. See text for details. Inside-out patches were bathed in either DVF or 1 mM Mg$^{2+}$-containing solution. C/A and I/O indicate cell-attached and inside-out. Application of PIP$_2$ and 1 mM La$^{3+}$ (La) in DVF solution are indicated. Time interval between data points (shown in circles) is 10 s. As described previously by us (Yeh et al., 2003, 2005), expression of TRPV5 in CHO cells is robust. We estimated that under our experimental condition each excised patch contains ~50–150 channels. PIP$_2$ reactivated run-down TRPV5 channels (i.e., currents recovered to >50% of the level before run-down) in 54 out of 62 recordings. We did not observe significant background currents in inside-out membranes of mock-transfected cells either before or after PIP$_2$ (inward [−100 mV] and outward [+100 mV] currents: −87 ± 32 pA and 117 ± 28 pA, respectively, before PIP$_2$ vs. −135 ± 41 pA and 156 ± 48 pA, respectively, after PIP$_2$; mean ± SEM, n = 6). (C) Ramp I-V curves of currents at indicated time points in B. Scale bars for X (40 ms) and Y axis (0.2 nA) are shown. Dotted line indicates 0 current. In the X axis, the time points corresponding to ramp potentials −100, 0, and +100 mV (from left to right) are indicated by three downward arrows, respectively. Inset shows I-V curves scaled up to the same peak current level.

Figure 1. Multiple effects of intracellular Mg$^{2+}$ on TRPV5. (A) Configuration and voltage protocol of recording. Voltage stimuli were applied every 10 s. Black and gray arrows indicate time points when inward and outward currents shown in Fig. 1 B were taken, respectively. (B) Intracellular Mg$^{2+}$ causes a fast reversible voltage-dependent block, a slow reversible inhibition, and an irreversible run-down of TRPV5 in inside-out membranes. See text for details. Inside-out patches were bathed in either DVF or 1 mM Mg$^{2+}$-containing solution. C/A and I/O indicate cell-attached and inside-out. Application of PIP$_2$ and 1 mM La$^{3+}$ (La) in DVF solution are indicated. Time interval between data points (shown in circles) is 10 s. As described previously by us (Yeh et al., 2003, 2005), expression of TRPV5 in CHO cells is robust. We estimated that under our experimental condition each excised patch contains ~50–150 channels. PIP$_2$ reactivated run-down TRPV5 channels (i.e., currents recovered to >50% of the level before run-down) in 54 out of 62 recordings. We did not observe significant background currents in inside-out membranes of mock-transfected cells either before or after PIP$_2$ (inward [−100 mV] and outward [+100 mV] currents: −87 ± 32 pA and 117 ± 28 pA, respectively, before PIP$_2$ vs. −135 ± 41 pA and 156 ± 48 pA, respectively, after PIP$_2$; mean ± SEM, n = 6). (C) Ramp I-V curves of currents at indicated time points in B. Scale bars for X (40 ms) and Y axis (0.2 nA) are shown. Dotted line indicates 0 current. In the X axis, the time points corresponding to ramp potentials −100, 0, and +100 mV (from left to right) are indicated by three downward arrows, respectively. Inset shows I-V curves scaled up to the same peak current level.

Effect Three: “Run-down” via Reduction of PIP$_2$. Membrane phospholipid PIP$_2$ regulates function of many ion channels and transporters (Hilgemann et al., 2001). In inside-out membranes, a prolonged exposure of the cytoplasmic face to Mg$^{2+}$ activates Mg$^{2+}$-dependent lipid phosphatases and depletes PIP$_2$ (Huang et al., 1998; Liu and Qin, 2005). Depletion of PIP$_2$ causes an irreversible loss of activity (“run-down”) of channels that are regulated by PIP$_2$ (Huang et al., 1998). For TRPV5, though each short exposure to Mg$^{2+}$ solution caused reversible inhibition, repetitive exposures to 1 mM Mg$^{2+}$ solutions led to progressive irreversible run-down of currents eventually (Fig. 1 B). Application of exogenous PIP$_2$ to inside-out membranes reactivated TRPV5 after its irreversible run-down by Mg$^{2+}$ (Fig. 1 B), suggesting that irreversible run-down of TRPV5 by intracellular Mg$^{2+}$ is due to loss of PIP$_2$ in the membrane and mechanistically different from the reversible voltage-dependent block and “slow inhibition” by Mg$^{2+}$. Activation of a Ca$^{2+}$-sensitive PLC by intracellular Ca$^{2+}$ depletes PIP$_2$ and leads to desensitization of TRPM8 channels (Rohács et al., 2005). We found that U-37122 (an inhibitor of PLC, 5–10 μM) did not prevent run-down of TRPV5 caused by prolonged exposure to Mg$^{2+}$ (unpublished data), supporting that Mg$^{2+}$-induced run-down is caused by activation of lipid phosphatase(s) rather than by activation of some Mg$^{2+}$-sensitive PLC.

A monoclonal antibody against PIP$_2$ (Fukami et al., 1988) interferes with PIP$_2$ regulation of ion channels presumably by sequestering PIP$_2$ in the membranes (Huang et al., 1998). Application of anti-PIP$_2$ antibody to the cytoplasmic face of inside-out membranes decreased TRPV5 currents (Fig. 2 A). Currents recovered partially after extensive washout of the antibody (Fig. 2 B). Polylysine, with its positively charged residues, has also been used to inhibit the activity of PIP$_2$-activated ion channels through competing PIP$_2$-channel inter-
trace 3 is due to free (ionized) Mg\(^{2+}\) by upward arrow. The characteristic voltage-dependent block in the time point corresponding to ramp potential 0 mV is indicated in Fig. 1A. Dotted line indicates 0 current. In the X axis, applied together with WMN (100 μM PIP2) and another solution receiving Mg-ATP were superimposed on one representative experiment showing long exposure of inside-out patches to Mg\(^{2+}\) causes run-down of TRPV5, which can be subsequently reactivated by Mg-ATP. // indicates 10 min break. The inhibition by Mg\(^{2+}\) shown here appears to be slower compared with that in Fig. 1B. This variability is likely due to differences in the configuration and accessibility of excised membranes to perfused bath solutions. Closed and open circles are currents at −100 and +100 mV, respectively. Similar results were observed in six separate experiments. In some experiments, Mg-ATP was applied together with WMN (10 μM). Gray triangles illustrate one such recording receiving Mg-ATP + WMN superimposed on one receiving the Mg-ATP alone. Similar results were observed in four separate experiments. Dimethyl sulfoxide (DMSO, vehicle for WMN, 0.5%) did not prevent reactivation by Mg-ATP (n = 4, not depicted). Of note is that wortmannin did not prevent recovery of currents from transient inhibition by Mg\(^{2+}\) (not depicted), indicating that the slow reversible inhibition is not due to regeneration of PIP2 from lipid kinases. (C) I-V curves of currents at indicated time points in B. In this experiment, voltage ramp from −100 to +100 mV was applied without a 40-ms step at −100 mV as indicated in Fig. 1A. Dotted line indicates 0 current. In the X axis, the time point corresponding to ramp potential 0 mV is indicated by upward arrow. The characteristic voltage-dependent block in trace 3 is due to free (ionized) Mg\(^{2+}\) (0.13 mM) present in the Mg-ATP solution.

Application of PIP2 to Inside-out Membranes Releases Mg\(^{2+}\)-induced Slow Reversible Inhibition

Activation of inward-rectifier K\(^{+}\) and other channels by PIP2 modulates the regulation of channels by other signaling pathways (Hilgemann et al., 2001). We examined whether PIP2 activation of TRPV5 alters its sensitivity to Mg\(^{2+}\)-induced slow inhibition. We examined dose–response relationships for the intracellular Mg\(^{2+}\)-induced slow reversible inhibition of TRPV5 in excised inside-out membranes before and after application of PIP2. As before, after reaching maximal currents in DVF solutions in inside-out membranes, TRPV5 currents were completely inhibited by a solution containing 1 mM Mg\(^{2+}\) (Fig. 3A). Currents recovered in DVF solutions. The concentration of Mg\(^{2+}\) for half-maximal inhibition (IC\(_{50}\)) of the inward currents of TRPV5 was 0.11 ± 0.03 mM (mean ± SEM, n = 7; Fig. 3C, closed circles). Application of exogenous PIP2 to inside-out membranes in DVF solutions increased TRPV5 currents by 2.4 ± 0.3-fold (n = 8; see Fig. 3A for representative experiment), indicating a submaximal concentration of PIP2 present in the excised membranes. The application of PIP2 reduced the sensitivity of TRPV5 to the slow reversible inhibition by intracellular Mg\(^{2+}\). After PIP2, the inward currents of TRPV5 were partially inhibited by 1 mM Mg\(^{2+}\) but completely inhibited by 10 mM Mg\(^{2+}\) (Fig. 3A). Currents recovered in the DVF solution (Fig. 3A). The IC\(_{50}\) for the slow inhibition by Mg\(^{2+}\) was increased by ~28-fold to 3.11 ± 0.25 mM (n = 8; Fig. 3C, open circles) after application of exogenous PIP2. The Mg\(^{2+}\)-induced voltage-dependent block remains distinguishable from the slow reversible inhibition even after application of exogenous PIP2. As shown by the scaled-up I-V curves (Fig. 3B), voltage-dependent block was complete in 1 mM Mg\(^{2+}\) in the presence of PIP2 (i.e., I-V curves at time point 2 and 3 are
superimposable, Fig. 3 B). Yet, Mg\(^{2+}\)/H\(^{+}\)-induced slow inhibition requires 10 mM Mg\(^{2+}\) over 30–40 s to reach its maximum (i.e., more inhibition occurred from time point 2 to time point 3 in Fig. 3 A).

**Figure 3.** PIP\(_2\) decreases the sensitivity to Mg\(^{2+}\)-induced slow inhibition but not the voltage-dependent block. (A) Application of PIP\(_2\) in inside-out patches increases TRPV5 currents and reduces the sensitivity to inhibition by Mg\(^{2+}\). Voltage ramp protocol is same as in Fig. 1. Holding potential between ramps is 0 mV. (B) Scaled-up I-V curves of currents at indicated time points in A. (C) Dose-response curves of inhibition by Mg\(^{2+}\) in inside-out patches before (no PIP\(_2\)) and after application of PIP\(_2\) (+PIP\(_2\)). IC\(_{50}\) was obtained by fitting relative inward currents (inward currents at −100 mV normalized to maximal currents; I/I\(_{\text{max}}\)) at different Mg\(^{2+}\) concentrations according to the equation I/I\(_{\text{max}}\) = IC\(_{50}\)/IC\(_{50}\) + [Mg\(^{2+}\)].

Effect of Intracellular Mg\(^{2+}\) on TRPV5 in Whole Cell
The concentration of PIP\(_2\) in the excised membranes is likely considerably lower than in the plasma membranes in vivo (Nasuhoglu et al., 2002). We further examined regulation of TRPV5 by intracellular Mg\(^{2+}\) using whole-cell recording (Fig. 4 A), which resembles physiological plasma membrane conditions better than the inside-out membranes. Strongly inwardly rectifying Na\(^{+}\) currents were observed immediately after establishment of whole-cell recording in TRPV5-expressing cells (time point 1 in Fig. 4 B and trace 1 in Fig. 4 C) but not in mock-transfected CHO cells (Fig. 4 D). Dialysis of the intracellular space with a DVF solution led to a large increase of whole-cell TRPV5 currents (Fig. 4, B and C).

As reported previously (Hoenderop et al., 2003; Yeh et al., 2003), whole-cell TRPV5 currents exhibited slight outward rectification at membrane potentials > +50 mV in the presence of extracellular EDTA (Fig. 4 C, trace 3). Perfusion of the intracellular space with a solution containing 1 mM Mg\(^{2+}\) caused a large reversible inhibition on whole-cell TRPV5 currents (Fig. 4, B and C). Unlike in the inside-out recordings, run-down of TRPV5 by Mg\(^{2+}\) is rather infrequent in whole-cell recordings (>90% of experiments show >80% recovery upon changing from Mg\(^{2+}\) to DVF solutions) despite an average longer exposure of the cytoplasmic face of plasma membrane to Mg\(^{2+}\). Thus, run-down of TRPV5 via activation of lipid phosphatases (as observed in excised inside-out patches) is likely not a physiological action of intracellular Mg\(^{2+}\) (as in the conditions of whole-cell recording). Intracellular Mg\(^{2+}\) at concentrations ranging from 0.1 to 3 mM caused a dose-dependent inhibition of whole-cell TRPV5 currents (Fig. 4 E). The IC\(_{50}\) for intracellular Mg\(^{2+}\) inhibition of TRPV5 channels in whole-cell recordings was 0.29 ± 0.02 mM (n = 11; Fig. 4 F). This IC\(_{50}\) is between that for inside-out membranes before PIP\(_2\) (IC\(_{50}\): 0.11 mM) and after PIP\(_2\) (IC\(_{50}\): 3.11 mM), suggesting that PIP\(_2\) content in whole cell membranes is intermediate between the latter two inside-out membrane conditions.

Regulation of ion channels including inward rectifier K\(^{+}\) and TRP channels by PIP\(_2\) involves direct interaction with positively charged amino acids in the channels (Hilgemann et al., 2001; Prescott and Julius, 2003; Rohács et al., 2005). Recently, Rohács et al. reported that arginine-606 of the rabbit TRPV5 (corresponding to arginine-599 of the rat TRPV5) is critical for interaction with PIP\(_2\) (Rohács et al., 2005). We found that R606Q mutant was more sensitive to inhibition by intracellular Mg\(^{2+}\) in whole-cell recordings (IC\(_{50}\): 0.03 ± 0.01 mM for R606Q [n = 4] vs. 0.29 ± 0.02 mM for wild type, P < 0.05; not depicted in Fig. 4). These results further support that PIP\(_2\) activation of TRPV5 releases intracellular Mg\(^{2+}\) inhibition of the channel.

Mg\(^{2+}\)-induced Slow Reversible Inhibition Is Voltage Dependent
To examine membrane voltage sensitivity, whole-cell TRPV5 currents were measured in intracellular DVF
solution and in a solution containing 0.3 mM Mg$^{2+}$ at different holding potentials. The fraction of TRPV5 currents in 0.3 mM intracellular Mg$^{2+}$ relative to maximal currents in DVF solution (I/Imax) increased with increasing membrane hyperpolarization (Fig. 5 A). Voltage sensitivity for the intracellular Mg$^{2+}$/H11001-induced slow reversible inhibition of TRPV5 is further supported by the finding in experiments using whole-cell recording that hyperpolarization of membrane holding potentials from 0 to $-50$ mV increased IC$_{50}$ for Mg$^{2+}$ from 0.29 to 0.79 mM (Fig. 5 B). Voltage dependency of inhibition by Mg$^{2+}$/H11001 can be viewed as Mg$^{2+}$/H11001 binds to inhibit the channel within the membrane electrical field. The fractional electrical distance of the binding site from the outer surface of the membrane ($\delta$) can be calculated according to equation $IC_{50}(V) = IC_{50}(0)\exp(26FV/RT)$ (Nilius et al., 2000), where IC$_{50}(V)$ and IC$_{50}(0)$ represent IC$_{50}$ at the test voltage $V$ and at 0 mV, respectively, and $R$, $T$, and $F$ have their usual meanings. Our results reveal that Mg$^{2+}$/H11001-induced slow inhibition of TRPV5 is owing to Mg$^{2+}$/H11001 binding inside the channel at $\delta = 0.26$.

**Time Constant Measurement Provides Further Support for Voltage Dependency for Mg$^{2+}$/H11001-induced Slow Inhibition and its Distinction from Voltage-dependent Block**

Up to now, our studies examine the regulation of TRPV5 by intracellular Mg$^{2+}$ by perfusion of solutions containing different concentration of Mg$^{2+}$. To avoid variability in the rate of perfusion and measure time constants for the Mg$^{2+}$/H11001-induced voltage-dependent block and the slow inhibition, we recorded whole-cell currents using voltage jump protocol at fixed intracellular Mg$^{2+}$ concentration (1 mM) (Fig. 6 A). Membrane potential was held at $-100$ mV. At this hyperpolarized potential, TRPV5 is fully open (not inhibited by intracellular Mg$^{2+}$) and conducts inward current (see Fig. 1). Stepping from $-100$ to $+100$ mV resulted in a fast (i.e., occurring in ms and reaching maximal inhibition in ~50 ms) time-dependent inhibition of the channel (shown as decrease of outward current in Fig. 6 A). This fast inhibition of currents (known as the Mg$^{2+}$/H11001-induced voltage-dependent block) represents the time course of binding of intracellular Mg$^{2+}$ into the pore and contributes to inward rectification of currents.

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**Figure 4. Regulation of TRPV5 by intracellular Mg$^{2+}$ in whole-cell recording.** (A) Alteration of intracellular Mg$^{2+}$ in whole-cell recording using intrapipette perfusion. (B) Intracellular perfusion of Mg$^{2+}$ (1 mM) in whole-cell recording inhibits TRPV5 reversibly. The Y axis is current density (nA/pF). Time bar for the X axis represents 4 min. (C) I-V curves of currents at indicated time points in B. (D) Time control of intracellular Mg$^{2+}$/H11001 perfusion in mock-transfected CHO cells (transfected with GFP plasmid alone). The Y axis is current density (pA/pF). Time bar for the X axis represents 4 min. (E) Dose-dependent inhibition of whole-cell TRPV5 currents by Mg$^{2+}$/H11001 (0–5 mM). Each experiment at a given Mg$^{2+}$/H11001 concentration was performed as in Fig. 4 B and inward currents were scaled to the same maximal (current at DVF solution; I/I$_{\text{max}} = 1$) and minimal level (current after La$^{3+}$/H11001; I/I$_{\text{max}} = 0$). (F) Dose–response curve of Mg$^{2+}$/H11001 inhibition in whole-cell recording (closed squares) vs. in inside-out patches before (closed circles; no PIP$_2$) and after PIP$_2$ (open circles; +PIP$_2$). IC$_{50}$ was calculated according to the equation $I/I_{\text{max}} = IC_{50}/IC_{50} + [Mg^{2+}]. I/I_{\text{max}} = $ relative currents normalized to maximal currents. The two curves for inside-out patches before and after PIP$_2$ are from Fig. 3 C.
through TRPV5 and TRPV6 channels (Hoenderop et al., 2003; Voets et al., 2003). As shown by others (Voets et al., 2003), the time course of voltage-dependent block could be fitted by two exponential terms. Fig. 6 B shows the two exponential time constants for voltage-dependent block at +25, +50, and +100 mV, respectively. Beyond the fast voltage-dependent block occurring within the first 50 ms, outward currents further decreased in a much slower time course (occurring in seconds and reaching minimal current level in ~60 s) (Fig. 6 C). The mean exponential time constant (τ) for the “Mg²⁺-induced slow inhibition” at +100 mV was 8.3 ± 1.1 s (mean ± SEM, n = 4) (Fig. 6 D). The on-rate constant (Kₘ) for slow inhibition by Mg²⁺ at +100 mV (Kₘ = 1/τ [Mg²⁺]) was estimated at 120 ± 14 s⁻¹M⁻¹ (mean ± SEM, n = 4). Outward currents remained relatively stable beyond the first 50 ms in the absence of intracellular Mg²⁺ (current level at 60 s = 91 ± 8% of that at 100 ms, n = 4; not depicted in Fig. 6). Consistent with voltage dependency of Mg²⁺-induced slow inhibition, time constant at +50 and +25 mV were 13.4 ± 1.2 s and 16.2 ± 0.4 s, respectively (Fig. 6 D, each P < 0.05 vs. time constant holding at +100 mV). These time constant values at +100, +50, and +25 mV are in agreement with the earlier observation that Mg²⁺-induced slow inhibition occurred over 30–40 s at 0 mV (see Fig. 1; taking into consideration reduced PIP2 content in inside-out patches and probably a few seconds of delay in delivery of Mg²⁺ by perfusion).

Mutation of Aspartate-542 Abolishes Mg²⁺-induced Slow Reversible Inhibition

Acidic amino acids located in solvent-accessible region between the second intracellular loop (2nd IC loop) and the proximal region of the intracellular COOH terminus (CT) of TRPV5 may be involved in the binding and slow reversible inhibition by intracellular Mg²⁺ (Fig. 7 A). We mutated seven acidic amino acids within
this region, one at a time, to neutral amino acid aspar- 
agine or glutamine and tested the sensitivity of each 
mutant to inhibition by Mg2+ in whole-cell recordings. 
We found that neutralization of aspartate-542 (D542N) 
in the putative pore region completely abolished the 
slow reversible inhibition of TRPV5 by intracellular 
Mg2+ (Fig. 7, B and C). In voltage jump experiment, 
currents through D542N mutant remained unchanged 
over 60 s at +100 mV (current level at 60 s 89 ± 15% 
of that at 100 ms, n = 3; not depicted in Fig. 7), con-
firming that aspartate-542 is critical for Mg2+-induced 
slow inhibition. Neutralization of other aspartate or 
glutamate residues in this region had no significant ef-
facts on the slow reversible inhibition of TRPV5 by 
Mg2+ (Fig. 7 D).

Aspartate-542 is believed to be a part of the selectivity 
filter of TRPV5 and a binding site for Mg2+-induced 
voltage-dependent block (Nilius et al., 2001; Hoen-
derop et al., 2003). Our findings indicate that Mg2+ 
binding to aspartate-542 not only causes voltage-depen-
tent block but also the slow inhibition. This conclusion 
is further supported by the fact that the fractional ele-
ctrical distance for the slow inhibition we measured in 
this study (δ = 0.26) is similar to that measured for volt-
age-dependent block (δ = 0.31) (Nilius et al., 2000). 

Reduction of PIP2 Enhances Mg2+-induced Slow Reversible 
Inhibition of TRPV5 without Affecting Mg2+ Binding 
To investigate the mechanism for PIP2 modulation of 
Mg2+-induced slow reversible inhibition of TRPV5, we 
tested whether PIP2 affects Mg2+ binding. The kinetics 
of Mg2+-induced slow inhibition and voltage-depen-
tent block were studied using voltage jump exper-
iments as in Fig. 6. Whole-cell recordings were per-
formed in the presence of intracellular Mg2+ (1 mM). 
The extracellular bath solution contained thrombin + 
wortmannin (WMN) or vehicle. WMN inhibits phos-
phoinositide 4-kinase (a rate-limiting enzyme for syn-
thesis of PIP2 from phosphatidylinositol and phosphati-
dylinositol-4-phosphate) to allow reduction of mem-
brane PIP2 following receptor-mediated hydrolysis of 
PPIP2 (Nakanishi et al., 1995). Fig. 8 A shows representa-
tive single exponential fit to current traces in response 

Figure 7. Identification of amino acid of TRPV5 involved 
in Mg2+-induced slow inhibition. (A) Membrane topology 
of amino acids between the fourth transmembrane (TM4) 
and the proximal COOH terminus (Ct) of 
TRPV5. Amino acids in the preselectivity filter region 
likely form a pore helix struc-

ture similar to that of KcsA 
(Dodier et al., 2004; Voets et 
 al., 2004b). The location of 
aming acid mutated is shown. 
(B) A representative experi-
ment of D542N mutant (in 
whole-cell recording) showing 
no inhibition by 1 mM Mg2+. 
(C) Dose–response curves of 
inhibition by Mg2+ for D542N 
mutant and wild-type (WT) 
TRPV5. (D) Average inhibition 
by Mg2+ for WT and each 
mutant (mean ± SEM, n = 
5–11 each). * indicates P < 
0.05 vs. WT.
hydrolysis induced by receptor activation of PLC on TRPV5 functioning as a Ca\(^{2+}\)/H\(^{+}\)-permeable channel. In the following cell-attached recordings, the pipette solution contained (in mM) 140 NaCl, 1 MgCl\(_2\), and 1 CaCl\(_2\). Because of the anomalous mole fraction behavior, TRPV5 conducts Ca\(^{2+}\)/H\(^{+}\) exclusively in this physiological solution (Vennekens et al., 2001). CHO cells express endogenous receptors for thrombin (Dickenson and Hill, 1997). Activation of PLC by endogenous thrombin receptors in CHO cells was verified by the release of Ca\(^{2+}\) from intracellular stores (unpublished data). Thrombin was applied to the patch membrane via intrapipette perfusion (Fig. 9 A). The bath solution contained vehicle or wortmannin.

In cell-attached recording, application of thrombin (Thr) in the absence of WMN decreased TRPV5 Ca\(^{2+}\)/H\(^{+}\) currents transiently (by 43 ± 8% in 90 s, n = 5, P < 0.05 vs. control without thrombin) (Fig. 9 A). Presumably due to desensitization of receptors and de novo synthesis of PIP\(_2\) via phosphoinositide 4-kinase, TRPV5 currents recovered at least partially over 3–4 min in the absence of wortmannin. In the presence of wortmannin, thrombin caused a persistent reduction of TRPV5 currents by 85 ± 5% in 90 s (n = 6, P < 0.05 vs. thrombin alone). As controls, TRPV5 currents did not change significantly without thrombin (decreased by 8 ± 7% over 90 s, n = 4; WMN alone in Fig. 9 A). The effects of PIP\(_2\) hydrolysis on dose-dependent inhibition by Mg\(^{2+}\) of TRPV5-mediated Ca\(^{2+}\) currents were examined by whole-cell recording (Fig. 9 B). The IC\(_{50}\) for Mg\(^{2+}\) inhibition of TRPV5-mediated Ca\(^{2+}\) currents were 0.61 ± 0.12 mM (n = 5) and 0.11 ± 0.08 (n = 5) for control cells (treated with vehicle) and cells treated with thrombin and wortmannin, respectively (P < 0.05). The concentration of the intracellular free Mg\(^{2+}\) in the cell-attached recording is estimated at 0.5 mM (Romani and Scarpa, 2000). These results support the hypothesis that an increase in sensitivity to intracellular Mg\(^{2+}\) is important for physiological regulation of TRPV5-mediated Ca\(^{2+}\) entry by PLC-activating hormones.

**DISCUSSION**

Many TRP channels are downstream of G protein-coupled receptors and PLC (for review see Clapham, 2003; Putney, 2003). Thus, TRP channels may be regulated by G protein-coupled receptors via inositol-1,4,5-trisphosphate and/or diacylglycerol, products of PIP\(_2\) breakdown catalyzed by PLC (Clapham, 2003; Putney, 2003). Several TRP channels are also directly regulated by PIP\(_2\) (Chuang et al., 2001; Hardie, et al., 2001; Runnels et al., 2002; Liu and Liman, 2003; Liu and Qin, 2005; Rohács et al., 2005). Therefore, an additional mechanism for PLC to control TRP channels is via reduction of PIP\(_2\) content (Chuang et al., 2001; Runnels et al., 2002; Liu and Qin, 2005). However, the reduction of PIP\(_2\) in the plasma membrane from physiological stimulation of PLC may not be sufficient to alter the activity of channels if the affinity of channels for PIP\(_2\) is relatively high (Kobrinsky et al., 2000). In the present study, we find that PIP\(_2\) activates TRPV5 and that activation of TRPV5 by PIP\(_2\) decreases the sensitivity of
TRPV5 to inhibition by intracellular Mg2⁺ (see working model in Fig. 10). Hydrolysis of PIP2 by receptor activation of PLC increases the sensitivity to the inhibition by Mg2⁺. These results provide a novel mechanism for regulation of TRP channels by PLC-activating hormones by increasing the sensitivity to intracellular Mg2⁺.

The cytosolic free Mg2⁺ concentration in most mammalian cells is between 0.5 and 1 mM (Romani and Scarpa, 2000). The IC₅₀ for the slow inhibition (0.79 mM) at -50 mV (near the physiological resting membrane potential) suggests that intracellular Mg2⁺ is an important physiological regulator of TRPV5 function.

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Figure 9. Effect of PIP2 hydrolysis on the Mg2⁺-induced slow inhibition. (A) Effect of thrombin plus/minus wortmannin on TRPV5 in cell-attached recording. The top left panel shows cell-attached configuration for recording TRPV5-mediated Ca2⁺ currents. See text for the pipette solution. Voltage stimuli (step to -100 mV for 40 ms and ramp from -100 to +100 mV ramp over 200 ms) were applied every 2 s from +50 mV holding potential. Holding at +50 mV prevents Ca2⁺ entry and the Ca2⁺-dependent inactivation. Thrombin (Thr, 1 U/ml) was applied to the patch membrane by intrapipette perfusion. WMN (10 μM) or vehicle (0.5% DMSO) was applied to the extracellular bath in cells expressing TRPV5 5–10 min before addition of thrombin. The bottom panel shows inward currents (at -100 mV) normalized to the initial current (I/Imax). The top right panel shows ramp I-V curves at indicated time points. (B) Dose–response curve of inhibition by Mg2⁺ before and after membrane PIP2 hydrolysis. Whole-cell TRPV5-mediated Ca2⁺ currents were recorded in a bath solution containing (in mM) 140 NaCl, 1 MgCl₂, 1 CaCl₂, and 10 HEPES (pH 7.4). Intracellular Mg2⁺ (0 to 3 mM) was altered by intrapipette perfusion. Cells expressing TRPV5 were incubated with a bath solution containing either vehicle (closed circles) or WMN + Thr (open circles) for 5–10 min before patching for whole-cell recording. Voltage protocol and holding potential are as in the cell-attached experiment.

Figure 10. A working model showing the relationships of three different effects of Mg2⁺ on TRPV5. Binding of intracellular Mg2⁺ to D542 of TRPV5 causes a fast (in ms time scale) voltage-dependent block. Prolonged binding to the same site by Mg2⁺ (e.g., 10 s) leads to slow inhibition, which is likely due to conformational changes of the channel. The slow inhibition is also voltage dependent. Membrane PIP2 stabilizes TRPV5 in open conformation. PIP2 interaction with TRPV5 does not affect Mg2⁺ binding but prevents Mg2⁺-induced conformational changes. Mg2⁺ causes run-down by activating lipid phosphatases. However, run-down is probably not a physiological effect of Mg2⁺ as it occurs predominantly in excised inside-out membranes. Hydrolysis of PIP2 by activating PLC-coupled receptors sensitizes TRPV5 to inhibition by intracellular Mg2⁺. * indicates that time constant for Mg2⁺-induced slow inhibition of TRPV5 is shorter in PIP2-reduced (e.g., 5 s) than in PIP2-repleted state (e.g., 10 s). For comparison, Mg-induced slow inhibition has also been reported for TRPV6, TRPM6, and TRPM7 (Nadler et al., 2001; Runnels et al., 2001; Voets et al., 2003, 2004a). Mg2⁺-induced voltage-dependent block contributes to inward rectification of many channels, including inward-rectifier K⁺ channels and TRPV5 and 6. Mg2⁺ would presumably cause run-down on all PIP2-regulated ion channels.
in vivo. The intracellular free Mg²⁺ concentration changes only slightly in response to physiological stimuli (Romani and Scarpa, 2000). However, the dramatic shift in Mg²⁺ sensitivity by PIP₂ will provide amplification for regulation of TRPV5 by intracellular Mg²⁺ and PLC-activating hormones. One such example is the inhibition of calcium transport in the distal renal tubules by prostaglandin E₂ through activation of phospholipase C via EP₁ receptors (Hoenderop et al., 2002).

Besides TRPV5 and TRPV6, intracellular Mg²⁺ has recently been recognized as an important regulator of TRPM6 and TRPM7 channels (Nadler et al., 2001; Rnannes et al., 2001; Voets et al., 2004a). TRPM7 (also known as magnesium-inhibited cation channel [MIC]; Kozak et al., 2002; Prakriya and Lewis, 2002) is a ubiquitous channel important for cellular uptake of magnesium and cell viability (Schmitz et al., 2003). Magnesium ions entering cells through TRPM7 feedback on the channel to regulate its activity (Schmitz et al., 2003). TRPM7 is also regulated by PIP₂ (Rnannes et al., 2002). Whether PIP₂ also modulates Mg²⁺ sensitivity for TRPM7 is unknown at present.

TRPV5 does not conduct Mg²⁺ in physiological conditions. However, TRPM6 (a close homology of TRPM7) is a Mg²⁺-permeable channel responsible for magnesium transport in epithelial tissues (Schlingmann et al., 2002; Walder et al., 2002; Voets et al., 2004a). Both TRPV5 and TRPM6 are present in the distal convoluted tubule (DCT) cells (Hoenderop et al., 2002; Schlingmann et al., 2002; Walder et al., 2002; Voets et al., 2004a), which are key cells responsible for transcellular reabsorption of magnesium and calcium in the kidney (Suki et al., 2000). Many diseases with reduced Mg²⁺ transport in the DCT exhibit increased Ca²⁺ transport. These diseases include genetic diseases such as familial hypomagnesemia with hypocalciuria and Gitelman’s disease (Warnock, 2002) and acquired diseases such as cis-platinum toxicity (Mavichak et al., 1988). Our present finding that intracellular Mg²⁺ negatively regulates TRPV5 raises the possibility that a decrease in Mg²⁺ influx through TRPM6 leads to an increase in Ca²⁺ influx through TRPV5/TRPV6 in DCT cells, providing a possible molecular explanation for this disease phenotype. It should be noted that patients with mutations of TRPM6 develop hypocalcemia (Schlingmann et al., 2002; Walder et al., 2002), rather than hypercalcemia as would be expected from an increase in Ca²⁺ reabsorption in DCT cells alone. Hypocalcemia in these patients is believed secondary to parathyroid failure from severe magnesium wasting and hypomagnesemia (Anast et al., 1972).

Intracellular Mg²⁺ regulates inward rectification of ion channels including inward rectifier K⁺ channels and TRPV5 and TRPV6 channels via voltage-dependent block (Lu and MacKinnon, 1994; Hoenderop et al., 2003; Voets et al., 2003). In the present study, we report that intracellular Mg²⁺ regulates TRPV5 additionally via a mechanism likely involving conformational change of the protein (Fig. 10). We refer to this additional inhibition by Mg²⁺ as slow reversible inhibition to distinguish from the faster voltage-dependent block. A similar dual regulation of TRPV6 by intracellular Mg²⁺ via voltage-dependent block and a slower inhibition has recently been reported (Voets et al., 2003).

Aspartate-542 of TRPV5 is a part of the selectivity filter critical for permeation of Ca²⁺ ions and a binding site for Mg²⁺-induced voltage-dependent block of Ca²⁺ and Na⁺ currents (Nilius et al., 2001; Hoenderop et al., 2003). Our findings suggest that binding of intracellular Mg²⁺ to aspartate-542 in the selectivity filter inhibits TRPV5 via at least two separate but interrelated mechanisms. The first mechanism is by voltage-dependent block. In this mechanism, Mg²⁺ ions bind and occlude the channel pore, preventing ion permeation. This inhibition by blocking of channel pore occurs as soon as Mg²⁺ enters and binds the selectivity filter driven by membrane depolarization and reverses quickly by membrane hyperpolarization. This is similar to voltage-dependent block of inwardly rectifying K⁺ channels and other channels by Mg²⁺, polyamines, and other charged blockers (Hille, 2001). Binding of Mg²⁺ to the selectivity filter of TRPV5 for a longer duration, however, leads to the second mechanism of inhibition: the slow reversible inhibition. The slow reversible inhibition develops over 30–40 s and is not reversed by transient unbinding of Mg²⁺ ions during intermittent membrane hyperpolarization from voltage ramp applied every 10 s. The exact mechanism for the slow inhibition remains unknown, but likely involves conformational change(s) of the channel protein.

Our results also shed light on the mechanism by which PIP₂ alters the sensitivity of TRPV5 to slow inhibition by Mg²⁺. PIP₂ activates many ion channels directly by affecting their structure (Hilgemann et al., 2001). We find that PIP₂ activates TRPV5 in the absence of intracellular Mg²⁺ (Fig. 1 B). PIP₂ also regulates D542N, a Mg²⁺ binding site mutant (unpublished data). Thus, activation of TRPV5 by PIP₂ does not require its regulation by intracellular Mg²⁺. However, activation of TRPV5 by PIP₂ decreases the sensitivity of TRPV5 for intracellular Mg²⁺-induced slow inhibition. PIP₂ does so without affecting Mg²⁺ binding to TRPV5 (Fig. 8 C). We suggest that PIP₂ desensitizes TRPV5 to Mg²⁺-induced slow inhibition by stabilizing channel in an open conformation and prevents the conformational change induced by binding of Mg²⁺.

It is known that ion binding may alter the structure of selectivity filter of ion channels. For example, occupancy of K⁺ ions in the selectivity filter slows the rate of C-type inactivation of voltage-gated K⁺ channels.
(Yellen, 1998). The crystal structure of the bacterial KcsA K⁺ channel reveals that presence of multiple K⁺ ions in the selectivity filter stabilizes the full open conformation (Zhou and MacKinnon, 2003). Our present study provides an example of conformational changes of ion channels caused by binding of a pore blocker.

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