Divalent cations activate TRPV1 through promoting conformational change of the extracellular region

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Divalent cations Mg2+ and Ba2+ selectively and directly potentiate transient receptor potential vanilloid type 1 heat activation by lowering the activation threshold into the room temperature range. We found that Mg2+ potentiates channel activation only from the extracellular side; on the intracellular side, Mg2+ inhibits channel current. By dividing the extracellularly accessible region of the channel protein into small segments and perturbing the structure of each segment with sequence replacement mutations, we observed that the S1–S2 linker, the S3–S4 linker, and the pore turret are all required for Mg2+ potentiation. Sequence replacements at these regions substantially reduced or eliminated Mg2+-induced activation at room temperature while sparing capsaicin activation. Heat activation was affected by many, but not all, of these structural alternations. These observations indicate that extracellular linkers and the turret may interact with each other. Site-directed fluorescence resonance energy transfer measurements further revealed that, like heat, Mg2+ also induces structural changes in the pore turret. Interestingly, turret movement induced by Mg2+ precedes channel activation, suggesting that Mg2+-induced conformational change in the extracellular region most likely serves as the cause of channel activation instead of a coincidental or accommodating structural adjustment.

INTRODUCTION

Mg2+ was the first of many divalent and multivalent cations (such as Ni2+, Gd3+, and polyamines) shown to potently activate transient receptor potential vanilloid type 1 (TRPV1) (Ahern et al., 2005; Riera et al., 2007). It is thought that the potentiation effect of Mg2+ on TRPV1 activation may contribute to pain under certain conditions; indeed, intraperitoneal infusion of high concentrations of MgSO4 elicits identifiable pain responses in mice (Gyires and Torma, 1984; Mogil et al., 1999) that are mediated by TRPV1 (Ahern et al., 2005). The effect of Mg2+ (as well as other cations such as Ni2+) on TRPV1 gating can be prevented by mutations to two extracellularly located protonation sites in the outer pore region, suggesting that H+ and divalent cations might affect the same gating machinery (Jordt et al., 2000; Ahern et al., 2005; Luebbert et al., 2010; Wang et al., 2010). Our study described in our companion paper (see Cao et al. in this issue) suggests that Mg2+ and another divalent cation, Ba2+, directly affect heat activation while influencing other activation pathways that are allosterically coupled to the C→O transition of the channel pore. Mg2+ lowers the heat activation threshold into the room temperature range, leading to channel activation. In addition, both Mg2+ and Ba2+ induce channel desensitization to heat but not to capsaicin, suggesting that the divalent cation effect on heat activation is likely limited to the earlier heat-induced conformational change before the opening of the activation gate.

Given the tight linkage between divalent cations and heat activation, it is of great interest to understand how divalent cations interact with TRPV1 protein. Finding the location of channel structures that mediate the divalent cation effect is particularly attractive because numerous regions throughout the channel protein have been implicated to play a role in heat activation (Fig. 1) (Zheng, 2013b). Although allosteric modeling can mathematically describe effects of a stimulus on channel activation and its coupling with other stimuli, it does not provide structural evidence on the location of acting site(s) or the structure(s) transmitting stimulation to the activation gate. Here, we describe experiments aiming at identifying the structures that mediate Mg2+ potentiation of TRPV1 activation.

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Abbreviations used in this paper: FM, fluorescein maleimide; FRET, fluorescence resonance energy transfer; TMRM, tetramethylrhodamine maleimide; TRPV1, transient receptor potential vanilloid type 1.
MATERIALS AND METHODS

cDNA constructs

All constructs used in this study were based on the mouse TRPV1. Six sequence replacement mutants were made at the extracellular regions: three at the S1–S2 linker, one at the S3–S4 linker, and two at the pore turret (as illustrated in Figs. 3, 5, and 7). These mutations were designed to perturb the targeted structures and test their potential involvements in activation gating by Mg2+. For the S1–S2 linker mutations, the first five, middle six, and last six amino acids of the segment 456RPVEGLPPYKLNNTVGD472 were replaced with GGGGS, GGGGGS, and GGGGGS, yielding mutant Rs1s2_1, Rs1s2_2, and Rs1s2_3, respectively. For the S3–S4 linker mutation, the segment 534HRKE557 was replaced with GGGG, yielding mutant Rs3s4. Two more mutants were generated in the pore turret region and were named differently from the previous Rs1s2_1, Rs1s2_2, and Rs1s2_3, respectively. For the S3–S4 linker mutations, the S1–S2 linker, the first five, middle six, and last six amino acids of the segment 513PPHPKRGSACRPGN626, was replaced with GGGGSGGGGSGGGGS. For the second mutant, termed R3, a longer turret segment, 605GKNNSLPVESPHKCRGSACRPGN626, was replaced with GGGGSGGGGSGGGGS. These are among a group of pore turret mutants that we have studied previously, and their functional properties including heat response are described in a recent report (Gai et al., 2012).

Site-directed fluorescence recordings

Structural changes in the turret and other extracellular regions were monitored with site-directed fluorescence recordings using a patch fluorometry method as described previously (Yang et al., 2010). There are two extracellularly accessible cysteines in mouse TRPV1, C617 and C622, both located in the turret region. For fluorescence resonance energy transfer (FRET) experiments from the turret, we removed the second cysteine with a point mutation, C622A, allowing sulfhydryl-reactive fluorophores to be attached to C617. For FRET experiments from the extracellular S1–S2 linker, we removed both turret cysteines with alanine mutations and then introduced a point mutation, N467C. Functional properties of these cysteine mutant channels are described in a previous report (Yang et al., 2010). Fluorescein maleimide (FM) and tetramethylrhodamine maleimide (TMRM) were used to irreversibly label the cysteine. Fluorescence labeling was performed, after the establishment of whole-cell patch recording, with an FM/TMRM concentration ratio of 1:1 (1–5 mM in total). The labeling is expected to result in 87.5% of all channels containing both FM and TMRM. Completion of labeling was monitored by patch recording from the time course of change in current amplitude caused by attachment of fluorophores to the cysteine residues (Zheng and Zagotta, 2000). Free fluorophore molecules were thoroughly washed out before fluorescence recording. As multiple FRET pairs may form among fluorophores on the four channel subunits, the measured FRET efficiency reflected the sum of all possible fluorophore couplings and thus did not directly correlate to any single FRET pair (Bykova et al., 2006; Cheng et al., 2007; Zheng, 2013a). FRET changes reflected the overall effect of conformational change at the fluorophore attachment sites on distances between various FRET pairs. Potential contribution of nonspecifically labeled fluorophores was assessed using cells expressing a “cysteine-less” TRPV1 mutant, in which both extracellularly accessible cysteines were mutated to alanine.

FRET between FM and TMRM was measured with the Spectra FRET method (Zheng et al., 2002; Cheng et al., 2007) from voltage-clamped HEK293 cells imaged with an inverted fluorescence microscope (TE2000-U; Nikon) using a 40× oil-immersion objective (NA 1.3). An argon laser (Spectra-Physics) was used to provide the excitation light, with the exposure time controlled by a shutter (Uniblitz; Vincent Associates) synchronized with the camera by software through an amplifier (PatchMaster; HEKA). Two filter cubes (Chroma Technology Corp.) contained the following elements (excitation filter, dichroic mirror, and emission filter): Cube I, Z488/20, Z488rdc, and HQ500LP; Cube II, Z514/10, Z514rdc, and HQ530LP. For Spectra FRET, two spectroscopic images were taken from each cell at each measurement time point, one with the FM excitation at 488 nm using Cube I, and the other with the TMRM excitation at 514 nm using Cube II. From these two images, the total emission spectrum and the TMRM emission spectrum, respectively, were constructed using the linescan function in MetaMorph to determine the fluorescence intensity values along the wavelength axis. Standard emission spectra were collected from cells labeled with only TMRM or FM and used to separate cross-contamination between FM and TMRM caused by spectra overlaps (Takanishi et al., 2006). Spectral measurements were performed with a spectrograph (Acton SpectraPro 215i; Roper Scientific) in conjunction with a CCD camera (Roper Cascade 128B; Roper Scientific). The sample rate for fluorescence measurements was 1.14 Hz. The rates of photobleaching FM and TMRM by the excitation light were quantified separately from fluorophores attached to the channel and used to correct for photobleaching during FRET experiments.

FRET was quantified from the enhancement of TMRM fluorescence emission caused by energy transfer (Zheng et al., 2002; Cheng et al., 2007) using a method as described previously (Yang et al., 2010), with the following modifications. To assess potential artificial effect of Mg2+ on fluorescence emission, Mg2+-dependent changes in fluorescence intensity were measured from cells labeled with either FM or TMRM only. After labeling, the cells were initially bathed in a Na+-containing solution. After the basal fluorescence level was measured, solutions containing different concentrations of Mg2+ were perfused onto the cells while the fluorescence intensity was monitored. For both FM and TMRM, higher Mg2+ concentrations increased the fluorescence intensity by very similar fractions. There was no significant difference in the amplitude of Mg2+-induced fluorescence intensity change between FM and TMRM (Mg2+ concentrations tested were 10, 30, 100, and 130 mM; P = 0.62–0.97; n = 6 each). Therefore, for FM/TMRM intensity ratio measurements, no correction was applied. For FRET efficiency quantification, differences in fluorescence intensity caused by Mg2+ were corrected to ensure accuracy. To do so, we estimated the RatioA0 value in the absence and presence of...
Mg\textsuperscript{2+} potentiates TRPV1 heat activation only from the extracellular side

Experiments described in our companion paper (Cao et al., 2013) were all conducted with extracellular Mg\textsuperscript{2+}. To test whether Mg\textsuperscript{2+} applied to the intracellular side also potentiates TRPV1 heat activation, we repeated these experiments in inside-out patches with various concentrations of Mg\textsuperscript{2+} in the bath. As shown in Fig. 2 A, unlike extracellular Mg\textsuperscript{2+}, intracellular Mg\textsuperscript{2+} failed to elicit a current response from patches that yielded a large current upon capsaicin challenge. Closer inspection showed that intracellular Mg\textsuperscript{2+} inhibited spontaneous channel activity (Fig. S1 A), yielding an overall effect of reducing the current amplitude (Fig. 2 B). We further observed that intracellular Mg\textsuperscript{2+} inhibited capsaicin-induced current (Fig. 2 C; see also Fig. S1 B), which can be largely but not completely explained by an inhibitory effect on single-channel conductance (as described below). Inhibition of capsaicin-induced current could be rapidly and completely reversed upon washing off Mg\textsuperscript{2+} (Fig. 2 C).

We have described in our companion paper (Cao et al., 2013) that extracellular Mg\textsuperscript{2+} inhibits single-channel conductance to partially cover up its potentiation effect on gating. Intracellular Mg\textsuperscript{2+} exhibited a very similar dose-dependent inhibition of single-channel conductance.

Online supplemental material
Fig. S1 shows the lack of potentiation effect by intracellular Mg\textsuperscript{2+}. Fig. S2 shows capsaicin activation of the extracellular linker mutant channels. The online supplemental material is available at http://www.jgp.org/cgi/content/full/jgp.201311024/DC1.
reducing the current by ∼50% at the 100-mM concentration (Fig. 2, D–F). Even after correcting for conductance inhibition, there is no evidence that intracellular Mg$$^{2+}$$ activated the channel. Because Mg$$^{2+}$$ effect on single-channel conductance could be seen from either side of the channel, it is most likely that conductance inhibition occurred because Mg$$^{2+}$$ ions have a slow permeation rate, leading to blockade of conductance to other permeant ions. Such a permeation block mechanism has been discussed in great detail to explain observations from a wide range of ion channels (Neyton and Miller, 1988; Piasta et al., 2011).

Because Mg$$^{2+}$$ potentiates TRPV1 only from the extracellular side but not the intracellular side, the channel structure(s) that mediates the potentiation effect should be exposed to the extracellular aqueous environment. To search for the underlying structure(s), we generated systematic sequence alterations in the S1–S2 linker, the S3–S4 linker, and the pore turret, and tested the resultant mutants with extracellular Mg$$^{2+}$$ using both patch-clamp and live-cell Ca$$^{2+}$$ imaging recordings. Results from these experiments are described next.

Mutations of the S1–S2 linker can eliminate potentiation effect of extracellular Mg$$^{2+}$$

To test for potential involvement of the S1–S2 linker in Mg$$^{2+}$$-induced activation, we first replaced the sequence between R456 and D472 with a (GGGGS)$$^3$$ sequence. However, this mutant construct failed to yield functional current when expressed in HEK293 cells. To circumvent this problem, we divided the S1–S2 linker into three segments and replaced each with a GnS sequence of equal length, where n represents the number of glycine residues (Fig. 3 A). All three constructs yielded functional channels. Resembling many pore turret mutants we studied previously (Yang et al., 2010; Cui et al., 2012), S1–S2 linker mutants exhibited well-preserved capsaicin activation. Accordingly, cells expressing these mutants yielded robust intracellular Ca$$^{2+}$$ signals upon capsaicin challenge (Fig. 3 B). An example dose–response relationship for Rs1s2_3 is shown in Fig. S2. However, although the wild-type channel yielded a strong fluorescence response to 130 mM of extracellular Mg$$^{2+}$$ that was on average 87.3 ± 2.7% ($$n$$ = 48) in amplitude compared with that elicited by 10 µM capsaicin, both Rs1s2_1 reducing the current by $$\leq$$ 50% at the 100-mM concentration (Fig. 2, D–F). Even after correcting for conductance inhibition, there is no evidence that intracellular Mg$$^{2+}$$ activated the channel. Because Mg$$^{2+}$$ effect on single-channel conductance could be seen from either side of the channel, it is most likely that conductance inhibition occurred because Mg$$^{2+}$$ ions have a slow permeation rate, leading to blockade of conductance to other permeant ions. Such a permeation block mechanism has been discussed in great detail to explain observations from a wide range of ion channels (Neyton and Miller, 1988; Piasta et al., 2011).

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Figure 3. Two out of three S1–S2 linker mutants exhibit a lack of Mg$$^{2+}$$ potentiation. (A) Diagram illustrating the design of mutations. (B) Representative Ca$$^{2+}$$ imaging traces. (C) Summary of the amplitude ratio between Mg$$^{2+}$$- and capsaicin-induced fluorescence changes in Ca$$^{2+}$$ imaging experiments. $n$ = 48 (wild type), 42 (Rs1s2_1), 55 (Rs1s2_2), and 89 (Rs1s2_3). (D) Representative current traces from Rs1s2_1 (top) and Rs1s2_2 (bottom) in response to 130 mM Mg$$^{2+}$$ and 10 µM capsaicin. (E) Summary of the amplitude ratio between 130 mM Mg$$^{2+}$$-induced current and 10 µM capsaicin-induced current. $n$ = 5 (wild type), 4 (Rs1s2_1), and 3 (Rs1s2_2). (F) Representative current trace (top) and comparison of the amplitude ratio (bottom) between Mg$$^{2+}$$- and capsaicin-induced currents for wild type (black bars) and Rs1s2_3 (blue bars). $n$ = 5 (wild type) and 3–6 (Rs1s2_3). *, $P$ < 0.05; ***, $P$ < 0.001; N.S., not significant.
and Rs1s2_2 failed to show a detectable response to Mg\textsuperscript{2+} (P < 0.001; n = 42 and 55, respectively) (Fig. 3, B and C). For Rs1s2_3, a robust response to Mg\textsuperscript{2+} was reliably detected, although the amplitude appeared to be slightly lower than that of the wild-type channel (P < 0.05; n = 89).

Patch-clamp recordings confirmed the observations from Ca\textsuperscript{2+} imaging experiments, showing a nearly complete absence of Mg\textsuperscript{2+} response from Rs1s2_1 and Rs1s2_2 but a strong response from Rs1s2_3 (Fig. 3, D–F). We also observed that, although a strong capsaicin response could be recorded from both Rs1s2_1 and Rs1s2_2 after the failed attempt with Mg\textsuperscript{2+}, the capsaicin response was nonetheless slow, which was particularly true for Rs1s2_2 (n = 4 and 3, respectively) (Fig. 3 D). Recovery from capsaicin activation also appeared to be much slower, allowing us to confirm Mg\textsuperscript{2+} inhibition of the conductance of these mutant channels during the recovery time course (Fig. 3 D). Furthermore, Mg\textsuperscript{2+}-induced current from Rs1s2_3 desensitized slowly in the presence of Mg\textsuperscript{2+} (Fig. 3 F), which closely resembled the desensitization behavior of the wild-type channel. In summary, although there are interesting differences in how each S1–S2 linker mutant behaved, the basic observation is that two of them lost the ability to respond to extracellular Mg\textsuperscript{2+}, whereas the third exhibited a mildly reduced response.

Impaired heat response of the S1–S2 linker mutants
As Mg\textsuperscript{2+} activation and heat activation in the wild-type channel are tightly coupled, we tested how each S1–S2 linker mutant responded to heat. Ca\textsuperscript{2+} imaging recordings demonstrated that, although heating to 40°C and higher could elicit a strong fluorescence response from both the wild-type channel and Rs1s2_3, the same stimulus failed to elicit a response from cells expressing Rs1s2_1 or Rs1s2_2 (even though the cells were responsive to capsaicin, confirming positive expression of functional channels) (Fig. 4, A and B). Again, the response of Rs1s2_3 to heat is similar but statistically smaller than that of the wild-type channel (P < 0.05; n = 52).

Results from patch-clamp recordings are consistent with those from Ca\textsuperscript{2+} imaging experiments. As shown in Fig. 4 (C and D), heating to temperatures substantially above the activation threshold of the wild-type channel failed to activate Rs1s2_1 and Rs1s2_2, whereas capsaicin activated these channels to high open probabilities.

![Figure 4](image_url)

**Figure 4.** Two out of three S1–S2 linker mutants do not respond to heat. (A) Representative Ca\textsuperscript{2+} imaging traces. (B) Summary of the amplitude ratios between heat- and capsaicin-induced responses. n = 59 (wild type), 37 (Rs1s2_1), 45 (Rs1s2_2), and 52 (Rs1s2_3). *, P < 0.05; ***, P < 0.001. (C and D) Representative single-channel traces recorded at +80 mV from Rs1s2_1 (C) and Rs1s2_2 (D) at room temperature (RT), elevated temperatures, and in the presence of 10 μM capsaicin. The recording shown in D was from a patch containing two channels. (E) Representative current trace recorded from Rs1s2_3 in response to heating and 3 μM capsaicin. (F) Current-temperature relationship of Rs1s2_3. Dotted lines represent linear fits to the leak and channel currents.
96 Mg2+ promotes extracellular conformational change

Mutation of the S3–S4 linker eliminates Mg2+ activation but spares heat activation

We next examined the Rs3s4 mutant (Fig. 5 A). Both Ca2+ imaging and patch-clamp recordings showed that this mutant lost Mg2+-induced activation but remained sensitive to capsaicin challenge (Fig. 5, B–E, and Fig. S2). Based on the close correlation between Mg2+ activation and heat activation observed from the S1–S2 linker mutants, we anticipated that Rs3s4 would also show a lack of

In contrast, Rs1s2_3 was robustly activated by heat (Fig. 4 E). The activation threshold temperature of Rs1s2_3 was estimated at 36.9 ± 0.7°C (n = 6), which is close to the 37.7 ± 0.3°C (n = 9) value for the wild-type channel (P > 0.05).

In summary of all results from the S1–S2 linker mutants, we found that the first two thirds of the linker is required to support both Mg2+ and heat activation, and the last one third appears to have a minor role in both activation forms.

Figure 5. Mutation of the S3–S4 linker eliminates Mg2+ response. (A) Diagram illustrating the design of mutation. (B) Representative Ca2+ imaging trace. (C) Summary of the amplitude ratio between Mg2+ and capsaicin-induced fluorescence responses for wild type (black bar; n = 48) and Rs3s4 (blue bar; n = 65). (D) Representative current trace recorded at +80 mV in response to 130 mM Mg2+ and 3 µM capsaicin. (E) Summary of amplitude ratio between Mg2+ and capsaicin-induced current for wild type (black bar; n = 5) and Rs3s4 (blue bar; n = 4). ***, P < 0.001.

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Figure 6. Rs3s4 mutant exhibits heat-induced current activation. (A) Representative Ca2+ imaging trace (top) upon heat (bottom) and capsaicin challenges. (B) Summary of amplitude ratio between heat- and capsaicin-induced currents for wild type (black bar; n = 59) and Rs3s4 (blue bar; n = 55). ***, P < 0.001. (C) Representative single-channel traces recorded at +80 mV at varying temperatures and in the presence of 3 µM capsaicin. (D) Relationship between open probability (P_o) and temperature. Open circles represent average P_o of individual sweeps, and the red trace represents average P_o over a 3.5°C range. (E) Representative single-channel traces demonstrating rapid deactivation transition upon voltage step from +80 to −80 mV.
heat sensitivity. Indeed, Ca\textsuperscript{2+} imaging experiments showed that heating up to near 50°C barely induced any fluorescence signal at the resting membrane potential from cells expressing Rs3s4 (Fig. 6, A and B). Surprisingly, patch-clamp recordings at +80 mV showed that this mutant channel could be activated by heat (Fig. 6 C), with the activation threshold temperature being apparently similar to that of the wild-type channel (Fig. 6 D). We also observed that, although the channel could be activated to high open probabilities, this was only observed at a depolarized voltage of +80 mV. Upon repolarizing to −80 mV, the heat-activated channel promptly closed (Fig. 6 E). This rapid deactivation behavior resembles that of the wild-type channel (see, for example, Voets et al., 2004). It thus appears that Rs3s4 remains heat sensitive. Reasons for the lack of a fluorescence signal are unclear. Because the voltage-dependent process has a substantial influence on channel activation when the heat activation pathway is potentiated, one possibility for the lack of a fluorescence response in live cells may be associated with the negative resting membrane potential under our experimental condition that might keep this mutant channel in the closed state. In summary, mutation at the S3–S4 linker eliminates Mg\textsuperscript{2+} activation but not heat activation. The fact that heat can activate Mg\textsuperscript{2+}-insensitive mutant channels indicates that the two gating modalities may be supported by only partially overlapping channel structures.

Mutations in the pore turret simultaneously affect Mg\textsuperscript{2+} and heat activations

We applied the same strategy to test potential involvement of the pore turret in Mg\textsuperscript{2+} activation (Fig. 7 A). The pore turret has been shown to participate in heat activation; several turret mutants exhibited diminished heat response (Yang et al., 2010; Cui et al., 2012). One such turret mutant, R3, was found to also exhibit a near complete absence of Mg\textsuperscript{2+}-induced fluorescence signal (Fig. 7, B and C). Patch-clamp recordings further showed that Mg\textsuperscript{2+} barely elicited any current from R3-expressing cells that responded strongly to capsaicin (Fig. 7, D–F). Similarly, Ba\textsuperscript{2+} also failed to activate R3 (Fig. 7, G–I). These findings were further supported by observations from a less severe turret mutant, R2. Like R3, R2 exhibited substantially reduced Mg\textsuperscript{2+}-sensitive fluorescence signals (Fig. 7, B and C). In addition, both Mg\textsuperscript{2+} and

Figure 7. Pore turret mutants exhibit diminished Mg\textsuperscript{2+} response. (A) Diagram illustrating the design of mutations. (B) Representative Ca\textsuperscript{2+} imaging traces. (C) Summary of amplitude ratio between Mg\textsuperscript{2+}- and capsaicin-induced fluorescence responses. n = 48 (wild type), 38 (R2), and 66 (R3). (D and E) Representative current trace of wild type (D) and R3 (E) challenged by Mg\textsuperscript{2+} and capsaicin. (F) Summary of amplitude ratio between Mg\textsuperscript{2+}- and capsaicin-induced currents. n = 5 (wild type), 9 (R2), and 5 (R3). (G and H) Representative current trace of wild type (G) and R3 (H) challenged by Ba\textsuperscript{2+} and capsaicin. (I) Summary of amplitude ratio between Ba\textsuperscript{2+}- and capsaicin-induced currents. n = 5 (wild type), 6 (R2), and 4 (R3). **, P < 0.01; ***, P < 0.001.
Ba\(^{2+}\) induced a much smaller current from R2 than the wild-type channel (Fig. 7, F and I). Clearly, the pore turret mutants that affected heat activation also significantly diminished divalent cation–induced activation.

**Extracellular Mg\(^{2+}\) promotes turret conformational change**

Heat activation of TRPV1 involves turret movement that can be directly observed from changes in FRET between fluorophores attached to the turret (Yang et al., 2010). If Mg\(^{2+}\) promotes heat activation, can Mg\(^{2+}\) induce turret movement in the absence of a temperature change? To answer this question, we performed patch fluorometry measurements (Fig. 8 A). FM and TMRM were attached to C617 in the middle of the turret (after removing another turret cysteine by a C622A mutation). Similar to the wild-type channels, fluorophore-labeled channels were reversibly potentiated by Mg\(^{2+}\) at room temperature, which can be seen by patch-clamp current recordings (Fig. 8 B, open circles; \(n = 14\)). The ratio between TMRM and FM fluorescence intensities was also increased by Mg\(^{2+}\), with the change being reversible upon removal of Mg\(^{2+}\) (Fig. 8 B, red circles). An increase in the TMRM/FM intensity ratio indicated an increase in FRET, suggesting that the fluorophores moved closer to each other, just like when heat was used to activate the channel (Yang et al., 2010). In addition, an “overshoot” in the TMRM/FM ratio was often observed at the end of the raising phase; upon removal of Mg\(^{2+}\), an increase in the TMRM/FM ratio was also often seen before the fluorescence decline phase (Fig. 8 B). As a negative control, the TMRM/FM ratio of untransfected cells was monitored under the same experimental conditions but was found to be completely insensitive to changes in Mg\(^{2+}\) concentration (Fig. 8 C). Thus, the observed changes in the TMRM/FM ratio at C617 indicate that Mg\(^{2+}\) induced a turret movement at room temperature, and the movement likely proceeded in multiple steps.

To further confirm that changes of the TMRM/FM ratio at the pore turret reflected Mg\(^{2+}\)-induced conformational changes, we repeated the patch fluorometry experiment with a mutant channel whose two turret cysteines were removed, and a new cysteine was introduced into the S1–S2 linker at N467, the first amino acid of the Rs\(_{1s2_3}\) mutation. Because the Rs\(_{1s2_3}\) mutant channel responded to Mg\(^{2+}\) similarly to the wild-type channel, it is likely that the structure near N467 does not participate in Mg\(^{2+}\)-induced structural changes. Indeed, we found that the application of Mg\(^{2+}\) failed to induce any detectable change in the TMRM/FM ratio, even though the labeled channels could be substantially activated by Mg\(^{2+}\) like the wild-type and Rs\(_{1s2_3}\) channels (Fig. 8 D; \(n = 6\)). Furthermore, when the experiment was repeated with channels missing both turret cysteines, we again observed Mg\(^{2+}\)-induced channel activation, but the background fluorescence did not change (Fig. 8 E; \(n = 3\)). These negative control experiments confirmed that the observed fluorescence changes at the pore turret are caused by Mg\(^{2+}\)-induced conformational change.

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**Figure 8.** Mg\(^{2+}\) induces pore turret movement.

(A) Bright-field and fluorescence images of a patch-clamped fluorescently labeled cell. Red dashed lines mark the position of the spectrograph input slit. Fluorescence signal from the area covered by the slit was collected by spectral imaging (right). (B; left) Simultaneous recordings of current (open circles) and TMRM/FM fluorescence intensity ratio (red circles) upon the application of 130 mM Mg\(^{2+}\). (Right) Spectral images and spectra taken at time points 1 and 2 labeled in the left panel. (C) Control experiments with untransfected cells labeled with FM or TMRM (\(n = 6\) each). No change in relative fluorescence intensity could be detected when various concentrations of Mg\(^{2+}\) were added. (D) Fluorescence signal recorded at N467C (which is close to the end of the S1–S2 linker) did not change upon Mg\(^{2+}\)-induced activation. (E) Background fluorescence did not change upon Mg\(^{2+}\)-induced activation of mutant channel without any extracellularly accessible cysteine.
regions are involved in the activation conformational change remains to be determined. The lack of a fluorescence change at the S1–S2 linker site (N467C) suggests that perhaps some of the peripheral regions might provide structural supports but do not directly participate in the Mg$^{2+}$-induced conformational change. If this is the case, it is most likely that the extracellular linkers and the pore turret interact with each other. More importantly, the mutational effects on Mg$^{2+}$ potentiation at extracellular linkers and the direct evidence of turret movement from the FRET experiments suggest that Mg$^{2+}$ induces a substantial conformational rearrangement of the channel’s extracellular structures that leads to channel activation.

The sidedness of Mg$^{2+}$ potentiation effect

An important finding of this study is that Mg$^{2+}$ potentiates heat activation only from the extracellular side but not from the intracellular side. Given its small size and high solubility in water, Mg$^{2+}$ should be in direct contact with any channel structure that is exposed to the aqueous environment. Should an intracellular channel structure undergo a large conformational change during heat activation, Mg$^{2+}$-induced turret movement is less substantial than heat-induced movement

To compare Mg$^{2+}$-induced turret conformational change to that induced by heat, we used the Spectra FRET method to quantify changes in FRET efficiency caused by turret movement (Fig. 9). We found that the magnitude of Mg$^{2+}$-induced FRET change is smaller than that of heat-induced FRET change at the same turret position (Fig. 9 C), which is consistent with the observation that Mg$^{2+}$ only partially activated TRPV1 (see, for example, Fig. 3 of our companion paper, Cao et al., 2013). As anticipated from a separation of the capsaicin and heat activation pathways, capsaicin failed to induce any FRET change at the pore turret (Fig. 9 C; $n = 5$) where heat and Mg$^{2+}$ induced positive FRET changes. In agreement with the TMRM/FM ratio measurements, there was no change in FRET efficiency at N467C or from the background fluorescence (Fig. 9 C; $n = 3–6$).

**Discussion**

Our results demonstrate that most of the extracellular channel regions are required to support Mg$^{2+}$ potentiation of TRPV1 activation. Participation of the channel’s extracellular regions in activation gating is not surprising, as mutations and H$^+$ binding to many extracellular sites have been found to have strong and often specific gating effects (Jordt et al., 2000; Myers et al., 2008; Grandl et al., 2010; Boukalova et al., 2013), and we have shown previously that the pore turret undergoes a substantial and specific conformational change during heat activation (Yang et al., 2010; Cui et al., 2012). What came as a surprise is how widely spread the participating regions are for Mg$^{2+}$ potentiation. Whether all the participating regions are involved in the activation conformational change remains to be determined. The lack of a fluorescence change at the S1–S2 linker site (N467C) suggests that perhaps some of the peripheral regions might provide structural supports but do not directly participate in the Mg$^{2+}$-induced conformational change. If this is the case, it is most likely that the extracellular linkers and the pore turret interact with each other. More importantly, the mutational effects on Mg$^{2+}$ potentiation at extracellular linkers and the direct evidence of turret movement from the FRET experiments suggest that Mg$^{2+}$ induces a substantial conformational rearrangement of the channel’s extracellular structures that leads to channel activation.

**Figure 9.** Mg$^{2+}$ induced FRET changes at the pore turret. (A) Mg$^{2+}$ did not affect RatioA$_0$ of TMRM-labeled cells (left) but increased RatioA of FM/TMRM-labeled cells (right), indicating an increase in FRET efficiency. (B) Statistical analysis of Mg$^{2+}$ effects on RatioA$_0$ (left) and RatioA (right). Dotted lines on the right link measurements from the same cells; RatioA$_0$ measurements were not paired. (C) Summary of Mg$^{2+}$-induced changes in FRET efficiency, which is defined as the FRET efficiency values in the presence of 100 mM Mg$^{2+}$ subtracted by the FRET efficiency value in the presence of Na$^+$. *, $P < 0.05$; **, $P < 0.01$. $n = 3–6$. 

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activation, having high millimolar concentrations of Mg$^{2+}$ in the surrounding solution is expected to exert an energetic influence on the stability of these conformations (Fersht, 1998), which is likely to be reflected by changes in channel activation. An excellent demonstration of this effect is that changes in ionic strength of the bath solution substantially affect voltage-sensor movement via the solvent-accessible cavities near S4 (Isals and Sigworth, 2001). Because the “heat sensor” of TRPV1 is expected to undergo a large conformational transition upon temperature change (Clapham and Miller, 2011), should it be exposed to the intracellular side of the channel, it would likely allow intracellularly applied Mg$^{2+}$ to affect heat activation. Therefore, our results suggest that heat-induced conformational changes to intracellular channel structures might be either minimal or limited to mostly buried protein structures.

Structural considerations of Mg$^{2+}$-induced TRPV1 activation

The structural architecture of TRPV1 is generally assumed to resemble that of the voltage-gated potassium channels, for which crystallographic studies revealed that the four turrets pose around the pore, the S1–S2 linkers locate near the turrets (with the S1 end pushing against the turret), and the S3–S4 linkers locate peripherally (Long et al., 2005, 2007) (Fig. 10 A). A low resolution cryo-EM study supports the anticipated structural similarity between TRPV1 and voltage-gated potassium channels (Moiseenkova-Bell et al., 2008). In potassium channels, the direct contact between the top of S1 and the pore region plays a critical role in voltage-dependent gating, perhaps by providing support for the voltage sensor to exert force onto the activation gate through the S4–S5 linker (Lee et al., 2009). Here, we observed in TRPV1 that the first two thirds of the S1–S2 linker (presumably in close proximity to the pore turret) and the turret itself are required to support Mg$^{2+}$ and heat activation, whereas mutations in the likely peripherally located S3–S4 linker and the S2 end of the S1–S2 linker only affected Mg$^{2+}$ activation or had minor effects on gating, respectively. These results agree with the existence of an overall structural similarity between TRPV1 and voltage-gated potassium channels (Fig. 10 B).

Observations described here and in our companion paper (Cao et al., 2013) indicate several gating steps during TRPV1 activation that are summarized in a schematic diagram shown in Fig. 10 C. Divalent cations cause a conformational rearrangement specifically to extracellular structures. This conformational rearrangement can be directly observed from the turret region with site-specific fluorophore labels, and can be disrupted by mutations to the turret as well as extracellular linkers between transmembrane helices. At least for the pore turret, the conformational change precedes pore opening, suggesting a likely cause–effect relationship. If

Figure 10. Divalent cations activate TRPV1 through conformational changes of the extracellular region. (A) Crystal structure of the Kv1.2-Kv2.1 chimera (Protein Data Bank accession no. 2R9R), with the pore turret, S1–S2 linker, and S3–S4 linker highlighted in surface-plot mode in different colors. The red circle highlights the S1–turret interaction that supports voltage-dependent gating. (B) Rosetta-generated structural model for TRPV1. (C) Schematic diagram summarizing the gating process of TRPV1 by divalent cations, heat, and capsaicin. Mg$^{2+}$/Ba$^{2+}$ and heat work from the extracellular side to induce a conformational rearrangement of channel structures exposed to the aqueous environment, which is coupled to pore opening (first horizontal transition). In the presence of Mg$^{2+}$/Ba$^{2+}$ or heat, the extracellular structure settles down over time to a more stable conformation that does not support the open pore conformation, leading to desensitization (second horizontal transition). Capsaicin binding to the intracellular S2–S3 linker region also leads to pore opening. This can happen to resting channels (left vertical transition) as well as to channels already desensitized to divalent cations (right vertical transition).
the stimulation by divalent cations or heat lasts long enough, a Ca\(^{2+}\)-independent conformational arrangement occurs, causing the channel to be desensitized to both divalent cations and heat. Although little is known about the nature of the desensitization process, we speculate that it is not caused by the closure of the activation gate (because capsaicin can still activate the desensitized channel). Instead, it may occur to the same channel structures that are affected by heat and divalent cations and are coupled to the activation gate. Capsaicin-induced activation is largely isolated from these extracellular processes and can proceed from either the resting state or the desensitized state. Details of these gating steps remain to be elucidated in future investigation of the heat activation mechanism.

**How do divalent cations potentiate TRPV1 heat activation?**

Although the study described in this and in our companion paper (Cao et al., 2013) identifies the pathway and channel structures that support the gating effects of divalent cations on TRPV1, the physical process underlying divalent cation–induced activation remains unclear. Several possibilities are considered briefly here. Based on observations that removing two extracellularly exposed proton-binding sites can eliminate the gating effect of many cations such as Mg\(^{2+}\), Ni\(^{2+}\), Ca\(^{2+}\), Gd\(^{3+}\), and polyamines (Ahern et al., 2005, 2006; Tousova et al., 2005; Ohta et al., 2008; Luebbert et al., 2010), it has been generally assumed that these cations bind to the same two sites. Proton binding to TRPV1 exerts a strong potentiation effect on gating that brings the open probability to apparently quite high levels (Tominaga et al., 1996). The gating effect is voltage dependent and substantially slow down channel activation kinetics (Terlau et al., 1998; Jordt et al., 2000). The two major proton-binding sites, E601 and E649 (corresponding to E600 and E648, originally identified in rat TRPV1) (Jordt et al., 2000), are both in the outer pore region (see Fig. 1). E601 is located at the junction between the S5 transmembrane domain and the pore turret, and E649 is located at the outer mouth of the pore near the selectivity filter. It can be envisioned that the binding of positive charges (H\(^{+}\), Mg\(^{2+}\), or other cations) at these positions may exert a substantial influence on the conformation of the channel pore or the surrounding outer pore structures such as the turret. In this scenario, the peripheral extracellular structures might play a supporting but nonetheless necessary role similar to what S1 does in potassium channels.

There are, however, noticeable differences between proton and divalent cations in potentiating TRPV1 activation. First, in all the mutant channels we tested, the two proton-binding sites remained unaltered. So to fit results from this study to the proton-binding site hypothesis, one has to assume that mutations in the broad extracellular regions altered the structure at the proton-binding sites so that Mg\(^{2+}\) can no longer bind there, or its binding fails to exert the same gating effect. Second, although the binding of H\(^{+}\) to the two outer pore sites occurs at micromolar concentrations (pH 6), we observed that it takes at least 10 mM Mg\(^{2+}\) to generate a detectable gating effect. In addition, no evidence for saturation of the potentiation effect was observed even at a concentration of 130 mM. Previous studies with Mg\(^{2+}\) (Ahern et al., 2005; Riera et al., 2007; Wang et al., 2010), Ca\(^{2+}\) and Sr\(^{2+}\) (Ahern et al., 2005), and Ni\(^{2+}\) (Luebbert et al., 2010) all yielded similar results. Only 30 µM Gd\(^{3+}\) and 500 µM spermine exhibited detectable gating effects at submillimolar concentrations at room temperature (Tousova et al., 2005; Ahern et al., 2006). Nonetheless, potentiation by polyamines does not always require the proton-binding sites; in fact, the E600Q mutation in rat TRPV1 enhanced the potentiation effect (Ahern et al., 2006). Gd\(^{3+}\) appears to indeed bind to the proton-binding sites, as it competes with H\(^{+}\) (Tousova et al., 2005).

Interaction between divalent cations and ion channels has been observed previously in many channel types. In most cases, Mg\(^{2+}\) of physiological concentrations (low millimolar) enters the channel pore to block ion flux, causing current rectification of the inward-rectifying potassium channels (Horie et al., 1987; Ciani and Ribaut, 1988) and current inhibition of the NMDA receptors (Nowak et al., 1984) and calcium channels (Lansman et al., 1986). In this study, we observed a similar conductance inhibition by Mg\(^{2+}\) in TRPV1. Mg\(^{2+}\) is found to also interact with the gating machinery of many channel types. In the case of ether-à-go-go channels, Mg\(^{2+}\) can substantially slow down channel activation kinetics (Terlau et al., 1996). The gating effect is voltage dependent and interestingly also involves competition between Mg\(^{2+}\) and H\(^{+}\) for common binding sites, which were later identified to be aspartate residues in the S2 and S3 segments (Silverman et al., 2000; Tang et al., 2000). Mg\(^{2+}\) can exert a substantial gating effect on ether-à-go-go channel gating at submillimolar concentrations. Because TRPV1 protein contains many charged (both positive and negative) residues that are likely exposed to the extracellular side (including the two H\(^{+}\)-binding glutamates), it is possible that Mg\(^{2+}\) can potentially interact with them to exert a gating effect.

As mentioned earlier, ionic strength has been shown to be able to exert a strong influence on protein stability and conformational change. Given that divalent cations and other molecules with varying chemical natures generally exert similar potentiation effects that are observable only at 10–100 mM concentrations (Riera et al., 2007; Ohta et al., 2008), it is also possible that they may potentiate heat activation not by binding to specific channel sites. Instead, high concentrations of divalent cations in solution may have a global (or perhaps localized) effect on channel protein stability. Through this nonspecific interaction, Mg\(^{2+}\) and Ba\(^{2+}\) contribute a favorable energetic effect on heat activation. The observation that widespread extracellular structures mediate the gating effect of divalent cations is consistent with...
this view. In a general sense, the process induced by high concentrations of divalent cations is not unlike protein denaturation induced by molar concentrations of urea. However, it is unlikely and unnecessary that complete denaturing of any part of the channel protein would occur during Mg\(^{2+}\)- and Ba\(^{2+}\)-induced activation. Instead, the gating machinery of TRPV1 shifts to a conformation that is normally unfavorable but becomes favorable at higher temperatures or in the presence of extracellular divalent cations.

In summary, widely spread extracellular structures are required for divalent cations to promote TRPV1 heat activation. Intracellular structures, however, are not involved in the potentiation process. As Mg\(^{2+}\)-induced turret movement precedes channel activation, it seems most likely that extracellular conformational rearrangement is coupled to the activation machinery and serves as the cause of channel activation rather than a coincidental or accommodating structural adjustment.

We are grateful to our laboratory members for assistance and helpful discussion throughout the course of the study, and Drs. Jon Sack and Vladimir Yarov-Yarovoy for critical reading of the manuscript.

This work was supported by a grant from the National Institutes of Health (R01 NS072577) and a UC Davis Health System Research Award (to J. Zheng); grants from the Ministry of Education of China 111 Project (B07001), the Ministry of Science and Technology of China (2013CB331302), and the National Science Foundation of China (81221002 to K.W. Wang); an Australian National Health and Medical Research Council fellowship (to L. Ma); and an American Heart Association predoctoral fellowship (to F. Yang).

The authors have no conflicting financial interests.

Sharona E. Gordon served as editor.

Submitted: 9 May 2013
Accepted: 22 November 2013

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