Divalent Heavy Metal Cations Block the TRPV1 Ca\(^{2+}\) Channel

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Abstract Transient receptor potential vanilloid 1 (TRPV1) is a non-selective cation channel involved in pain sensation and in a wide range of non-pain-related physiological and pathological conditions. The aim of the present study was to explore the effects of selected heavy metal cations on the function of TRPV1. The cations ranked in the following sequence of pore-blocking activity: Co\(^{2+}\) (half-maximal inhibitory concentration (IC\(_{50}\))=13 μM)>Cd\(^{2+}\) (IC\(_{50}\)=38 μM)>Ni\(^{2+}\) (IC\(_{50}\)=62 μM)>Cu\(^{2+}\) (IC\(_{50}\)=200 μM). Zn\(^{2+}\) proved to be a weak (IC\(_{50}\)=27 μM) and only partial inhibitor of the channel function, whereas Mg\(^{2+}\), Mn\(^{2+}\) and La\(^{3+}\) did not exhibit any substantial effect. Co\(^{2+}\), the most potent channel blocker, was able not only to compete with Ca\(^{2+}\) but also to pass with it through the open channel of TRPV1. In response to heat activation or vanilloid treatment, Co\(^{2+}\) accumulation was verified in TRPV1-transfected cell lines and in the TRPV1+ dorsal root ganglion neurons. The inhibitory effect was also demonstrated in vivo. Co\(^{2+}\) applied together with vanilloid agonists attenuated the nocifensive eye wipe response in mice. Different rat TRPV1 pore point mutants (Y627W, N628W, D646N and E651W) were created that can validate the binding site of previously used channel blockers in agonist-evoked \(^{45}\)Ca\(^{2+}\) influx assays in cells expressing TRPV1. The IC\(_{50}\) of Co\(^{2+}\) on these point mutants were determined to be reasonably comparable to those on the wild type, which suggests that divalent cations passing through the TRPV1 channel use the same negatively charged amino acids as Ca\(^{2+}\).

Keywords Heavy metals · Somatosensory system · Pain · Calcium channel · TRPV1 · Cobalt

Introduction

Small-diameter sensory neurons in the peripheral nervous system (PNS) express the transient receptor potential/vanilloid receptor subtype 1 (TRPV1). Four identical subunits of this protein form a functional Ca\(^{2+}\) channel. Similarly to other transient receptor potential channel members, the TRPV1 channel belongs in the large superfamily of cation channels with six transmembrane (TM) segments. Following agonist-induced channel opening, a pore loop between segments TM5 and TM6 serves as a cation filter and entry site [1].

When endovanilloids are produced under various inflammatory conditions around the C and Aδ afferents of these PNS neurons, TRPV1 transmits a specific pain sensation to the brain [2–4]. Besides endovanilloids such as anandamide [5, 6], TRPV1 is activated by acids (pH<6.3) and chemoirritants such as the exovanilloid capsaicin (CAPS) or resiniferatoxin, a naturally occurring, ultrapotent CAPS analogue with excellent specificity for TRPV1 [2, 7–9]. Moreover,
TRPV1 can integrate the effects of heat and vanilloids. The heat-sensing domain has been mapped to the C-terminal intracellular region [3, 9, 10]. The vanilloid binding site is localized between segments TM3 and TM4 of TRPV1 [5]. The pain signal, however, is generated by opening of the Ca\(^{2+}\) channel situated between domains TM5 and TM6 of four identical subunits [11, 12]. An acidic peptide motif in the pore loop region of TRPV1 (\textit{DXEXXXEXXD}) may serve as a docking site for positively charged ions and channel blockers [13]. However, our sequence comparison and previous in silico modeling efforts have not revealed any obvious homology to other divalent metal ion (M\(^{2+}\))-binding structures such as the \textit{EF-hand} (\textit{DXDXDGXXDXXE}) or the \textit{Excalibur} (\textit{DXDXDXXXXCE}) [13].

Various studies have demonstrated that positively charged molecules can act as TRPV1 receptor channel blockers by plugging the pore. Ruthenium Red (RuRed) (a well-known, but non-specific TRPV1 pore blocker) [11], R\(_4\)W\(_2\) (a positively charged hexapeptide) [14] and anti-calmodulins/antipsychotic tricyclics [13] have been shown to be able to bind to the \textit{DXEXXXEXXD} domain of TRPV1, whereby they block the movement of Ca\(^{2+}\) through the pore region.

We set out to assess the effects of various metal cations at different concentrations on the vanilloid - or heat-induced activity of the TRPV1 channel, focusing on the investigation of the most potent cations in vitro and in vivo. Moreover, in our experiments we aimed to shed light on the characteristics of the gating of the TRPV1 channel in order to improve the understanding of the structure and function of the TRPV1 pore region, which may lead to the development of potentially useful painkiller drugs that modulate the activity of this receptor.

**Materials and Methods**

**Reagents** Stock solutions (200 mM) of CoCl\(_2\), NiCl\(_2\), ZnSO\(_4\), CdCl\(_2\), CuSO\(_4\), Ca\(_2\)Cl\(_2\), CoCl\(_2\) and LaCl\(_3\) were dissolved in water and diluted as required to the working concentrations. To avoid the precipitation of insoluble La(OH)\(_3\) and La (CO\(_3\))\(_3\), the formation of radiocolloids and the loss of La\(^{3+}\) by adsorption to container surfaces, LaCl\(_3\) solution was prepared fresh daily in polyethylene vials [15]. RuRed and capsaicin (CapZ; Sigma-Aldrich, St. Louis, MO) were dissolved in DMSO. CAPS was dissolved initially as a stock solution of 3 mM in 95 % ethanol. The peptide R\(_4\)W\(_2\) was synthesized in our laboratories and then dissolved in water and used as a 25 mM stock solution. Amitriptyline (AMI), purchased from Sigma-Aldrich, was dissolved in water.

**Plasmids** The C-terminally epsilon-tagged rat TRPV1\(\varepsilon\) plasmid construct was prepared in the metallothionein (pMTH) plasmid vector as described earlier [9]. To avoid cell loss through the Ca\(^{2+}\)-excytotoxic mechanism that occurs when TRPV1 is overexpressed at 37 °C, only the basal activity of the pMTH promoter was used. The protein kinase C \(\varepsilon\) epitope tag allowed immunological detection of the TRPV1\(\varepsilon\) protein, as earlier described [16]. Mutants Y627W, N628W, D646N and E651W were kindly given by Dr. K. J. Swartz (National Institutes of Health, Bethesda, MD 20892, USA) [17] and subcloned into an EF-promoter-driven green fluorescent protein (EGFP)-tagging plasmid vector. The EGFP tag was used for visual determination of the transfection rate by flow cytofluorometry with a FACS-Calibur instrument (Becton Dickinson, San Jose, CA, USA).

**Cell Lines Expressing TRPV1 Ectopically** The HaCaT keratinocyte cell line was a kind gift of Prof. B. Farkas (Department of Dermatology, University of Cologne, Federal Republic of Germany) [18]. The COS-7 (CRL-1651) and BALB/c-3T3 (CCL-163) cell lines were obtained from ATCC. The 3T3 and HaCaT cell lines permanently expressing the rat TRPV1 channel were prepared as described earlier [19]. COS7 cells were transiently transfected with plasmid containing the sequence of the Y627W, N628W, D646N or E651W TRPV1 mutants or the wild-type TRPV1 channel, by using the Fugene transfection reagent (Roche, Mannheim, Germany). The transfection efficacy was determined by flow cytofluorometry.

**Primary Dorsal Root Ganglion Cultures** Were prepared from E16 embryonic rats as reported earlier [9]. Briefly, dorsal root ganglia (DRGs) were dissected and then processed in Hank’s balanced salt buffer until plated in Dulbecco’s Modified Eagle Medium (DMEM). The DMEM contained 20 mM HEPES, pH 7.4, 7.5 % foetal bovine serum, 7.5 % horse serum, 5 mg/ml uridine supplemented with 2 mg/ml 5-fluoro-2'-deoxyuridine and 40 ng/ml nerve growth factor to inhibit cell division and to promote the differentiation of long neuronal processes, respectively. Cells were seeded on 25 mm glass coverslips.

**Cobalt Histochemistry** Rat DRG cells attached to the coverslips were washed in buffer A (in millimolars: NaCl, 57.5; KCl, 5; MgCl\(_2\), 2; HEPES, 10; glucose, 12; sucrose, 139; pH 7.4) for 2 min, and then incubated at 37 °C for 10 min in Co\(^{2+}\)-uptake solution (buffer A+5 mM CoCl\(_2\)) containing 20 \(\mu\)M CAPS. High (20 \(\mu\)M) capsaicin concentration is used in order to obtain a robust and easily detectable Co\(^{2+}\) signal. Following a brief wash in buffer A, the water-soluble Co\(^{2+}\) taken up by the cells was precipitated with 0.12 % ammonium polysulphide (Sigma-Aldrich) in buffer A, which resulted in the formation of dark, water-insoluble CoS in TRPV1+ cells. Cells were fixed in 4 % formaldehyde and mounted on glass slides, using Kaiser’s glycerol.
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Statistical Analysis One-way ANOVA followed by Turkey’s post-tests was performed with GraphPad Prism version 3.01 software (GraphPad Software, Inc. San Diego, CA, USA).

Results

In the ⁴⁵Ca²⁺ uptake assay, the EC₅₀ of CAPS for wild-type TRPV1 was determined to be 0.0860 μM. Approximately 1 μM CAPS caused the full activation (EC₁₀₀) of TRPV1 at pH 7.5. Channel blocker-screening assays were therefore carried out with 2 μM CAPS (an excess amount of agonist), which does not cause Ca²⁺ cytotoxicity during the 10-min incubation period. The interactions of the metal ions with TRPV1 were studied by using a vanilloid-induced ⁴⁵Ca²⁺ uptake assay. Experiments were carried out on the TRPV1/ HaCaT permanent indicator cell line. Channel opening was induced by CAPS in the presence of progressively increasing M₂⁺ concentrations in the uptake solution. Incubation of the cells in uptake solutions containing both ⁴⁵Ca²⁺ and Mg²⁺, Mn²⁺ or La³⁺ (data not shown) resulted in little or no effect, even at the highest concentration (4 mM), whereas Zn²⁺ proved to be a weak [half-maximal inhibitory concentrations (IC₅₀)=27 μM] and only partial inhibitor of the 2 μM CAPS-induced ⁴⁵Ca²⁺ uptake. The other cations effectively blocked the vanilloid-induced Ca²⁺ entry into TRPV1/HaCaT cells, with the following sequence of potency: Co²⁺ (IC₅₀=13 μM)>Cd²⁺ (IC₅₀=38 μM)>Ni²⁺ (IC₅₀=62 μM)>Cu²⁺ (IC₅₀=200 μM; Fig. 1a).

To assess the effect of Co²⁺, the most potent TRPV1 inhibitor, on the heat-activated TRPV1 channels, the activity of TRPV1 was investigated in the presence either of 2 μM CAPS alone or of 2 μM CAPS+250 μM Co²⁺, at both 37 and 42 °C. The negative control did not contain CAPS. In this assay, high temperature activated the TRPV1 channels and also increased the CAPS-evoked ⁴⁵Ca²⁺ influx. Co²⁺ reduced both the heat and CAPS-induced ⁴⁵Ca²⁺ influx (Fig. 1b).

To compare the potency of Co²⁺ with those of the other positively charged channel blockers, we measured the IC₅₀ values of RuRed, AMI and R₄W₂, which are known to have a docking site in the pore loop of TRPV1. The inhibitor potentials of these pore blockers were measured via the CAPS-induced ⁴⁵Ca²⁺ uptake. All of them inhibited CAPS-activated TRPV1, with the following IC₅₀ values: RuRed=1 μM, AMI=20 μM and R₄W₂=100 μM (Fig. 1c).

For a better understanding of the inhibition kinetics of Co²⁺ on TRPV1, increasing concentrations of both Co²⁺ and CAPS were applied in the vanilloid-induced ⁴⁵Ca²⁺-uptake assays. The Ca²⁺ uptake of TRPV1/HaCaT cells was inhibited by the simultaneous presence of Co²⁺ in a dose-dependent manner. However, increasing concentrations of Co²⁺ decreased only the maximal response of efficacy.
of $Ca^{2+}$ entry; the affinity of CAPS for TRPV1 did not change. The inflection point in the CAPS dose–response curves in each of the $Co^{2+}$ co-incubation studies was found at $\sim 0.08 \mu M$ (i.e. EC$_{50}$), independently of the $Co^{2+}$ concentration. The $Co^{2+}$ inhibition patterns unequivocally indicated channel blocking kinetics (Fig. 2a).

By varying the concentrations of $Co^{2+}$ and $Ca^{2+}$ and measuring the radioactive $^{45}Ca^{2+}$ influx, we assessed whether there was a competition between $Co^{2+}$ and $Ca^{2+}$. The effect of dilution on the amount of accumulated $^{45}Ca^{2+}$ did not appear at extracellular cold $Ca^{2+}$ concentrations below 1 mM (Fig. 1a), indicating that TRPV1+ cells accumulate $Ca^{2+}$ very effectively from the extracellular space and collect them putatively into ER or mitochondria. Increasing cold $Ca^{2+}$ concentration decreased the inhibitory effect of $Co^{2+}$ (IC$_{50}$ values in the presence of 0, 15.625, 31.25, 62.5 and 125 $\mu M$ cold $Ca^{2+}$: 7.944, 51.22, 72.69, 79.09 and 189.1 $\mu M$, respectively), showing that the effect of $Co^{2+}$ on $Ca^{2+}$ entry mainly depends on the competition for entry sites. These results suggest that the $Co^{2+}$ entry through the TRPV1 channel is slower, and the $Co^{2+}$ displacing the $Ca^{2+}$ from the pore region of TRPV1 slows down or inhibits the $Ca^{2+}$ uptake (Fig. 2b).

The prolonged agonist stimulation of TRPV1 has been reported to result in an increased permeability to larger cations [22] or small molecules [23], due to conformational changes in the open state of the TRPV1. Thus, we analysed the kinetics of the channel-blocking activity of $Co^{2+}$ by employing different CAPS concentrations. An anticipated shift in the IC$_{50}$ of $Co^{2+}$ would be evidence supporting the idea that $Co^{2+}$ entry depends on the TRPV1 open stages. We indeed observed a shift in the IC$_{50}$ of $Co^{2+}$, which decreased with increasing CAPS concentration (Fig. 2c). Consequently, increasing agonist concentration enhances the blocking ability of $Co^{2+}$. To investigate this phenomenon, we plotted the IC$_{50}$ values as a function of CAPS concentration. Curve-fitting analysis confirmed a strong interrelationship between IC$_{50}$ and the CAPS dose applied (Fig. 2d), suggesting that the increased efficiency of inhibition correlates with the different open-state conformations of the TRPV1 channel.

We traced $Co^{2+}$ upon vanilloid induction in sensory neuron cultures prepared from DRGs of rat embryos. To test $Co^{2+}$-accumulation patterns, cells were co-incubated with 20 $\mu M$ CAPS in $Co^{2+}$-containing $Ca^{2+}$-uptake medium, and the $Co^{2+}$ was then localized by means of NH$_4$S histochemistry. These experiments revealed that $Co^{2+}$ not only competes with $Ca^{2+}$ but also enters into the cytosol of specific PNS sensory neurons. Functionally responsive vanilloid-sensitive neurons (i.e. TRPV1+) exhibited dark-brown $Co^{2+}$ precipitates inside the rounded neuronal bodies (Fig. 3e). As expected from previous studies, TRPV1 is endogenously expressed in approximately one third of the cultured neurons [24–27]. Without CAPS, no intracellular $Co^{2+}$ accumulation was observed (data not shown). Similar experiments were carried out on rTRPV1/HaCaT and rTRPV1/3T3 cell lines. The accumulation of $Co^{2+}$ was blocked by RuRed, a channel blocker of heat and vanilloid pain signalling. Moreover, the dose-dependent inhibition of the cellular entry of $Co^{2+}$ was determined by the co-application of 5 $\mu M$ CapZ, a long-known competitive
antagonist of pungent vanilloids. After analysis of the photographs of the cells with the ImageJ program, statistical analysis of the data further confirmed our findings: the mean gray values of CAPS-exposed, CAPS-free and CapZ-exposed cells proved to be significantly different (CAPS without Co2+, 72.05±12.38 (S.D.), n=146; CAPS+Co2+, 124.4±21.51 (S.D.), n=111; CAPS+Co2+ + CapZ, 76.92±22.21 (S.D.), n=100; P values of the t tests: CAPS without Co2+ vs. CAPS+Co2+, P<0.0001; CAPS+Co2+ vs. CAPS+Co2+ + CapZ, P<0.0001; CAPS without Co2+ vs. CAPS+Co2+ + CapZ, P=0.0290; Fig. 3). The gray values were measured on the negatives of the images: the darker the cells, the higher the gray values. Analysis of the pictures in Fig. 4, showing TRPV1-expressing HaCaT cells, resulted in the same outcome. Following ANOVA, the groups were compared by using t tests. Each t test except that involving CAPS without Co2+ vs. CAPS+Co2++100 μM CapZ indicated a significant difference between the pairs of groups (P<0.05; Fig. 4). No substantial staining could be observed on 3T3 cells (Fig. 4). Statistical analysis of the gray values of the cells indicated no significant darkening in the absence of TRPV1 in the cell membrane.

In order to rule out the possibility that Co2+ can enter the cells through VGCCs, 3T3 cells were challenged with 50 mM extracellular KCl. These cells did not show any VGCC activity: the high extracellular KCl concentration-induced depolarization that opens the VGCC channels did not cause 45Ca2+ accumulation in the 45Ca2+ -uptake assay.

Moreover, the VGCC blocker nisoldipine did not decrease the CAPS-induced TRPV1-mediated 45Ca2+ accumulation (data not shown). No Co2+ staining was observed in the presence of 50 mM extracellular KCl (Fig. 5F–j). ANOVA indicated no significant differences among the groups (P=0.9150). These results confirm that the CAPS-induced Ca2+ and Co2+ influx in TRPV1/3T3 cells is due exclusively to the TRPV1 channel activity.

Besides the in vitro demonstration of Co2+ antagonism, we further validated this Co2+ inhibition phenomenon in tests of eye wiping in response to pungent vanilloids [28]. Co2+ again decreased the frequency of vanilloid-evoked defending movements. Inhibition experiments with CapZ cross-validated and confirmed our earlier findings (Fig. 6a).

To validate that Co2+ inhibition is a consequence of competition with Ca2+ for M2+-chelating sites in the pore loop domain, we prepared several point mutants in this region of TRPV1, and determined the channel kinetics through 45Ca2+ -uptake experiments in 3T3 cells expressing the mutant channels. Some of these mutants had been partially characterized earlier in the context of spider venom channel inhibitors [17], and these residues proved to have an important role in the binding of various previously tested channel blockers (RuRed, etc.). Mutated sites are illustrated schematically in Fig. 6b.

The EC50 and EC100 values of mutant channels for CAPS were different from those of the wild type (D646N EC50=270 nM, E651W EC50=540 nM, N628W EC50=720 nM and Y627W EC50=820 nM), and the blocking effects of Co2+ and
Fig. 3 Co\textsuperscript{2+} histochemistry on the 3T3 cell line expressing TRPV1 ectopically. Cells were incubated for 10 min in buffer A containing a 20 μM CAPS+5 mM Co\textsuperscript{2+}; b 20 μM CAPS+5 mM Co\textsuperscript{2+}+5 μM CapZ; c 5 mM Co\textsuperscript{2+} without CAPS. The dark CoS precipitate indicates the presence of intracellular Co\textsuperscript{2+} that is blockable with CapZ, a competitive antagonist of pungent vanilloids. d Gray values of the 3T3 cell line expressing TRPV1 measured by means of ImageJ software. e Co\textsuperscript{2+} histochemistry in CAPS-sensitive rat DRG neurons. Scale bar=0.05 mm. These results confirm that Co\textsuperscript{2+} not only acts as a blocker but also enters the cell with Ca\textsuperscript{2+} through the TRPV1 channel.

Fig. 4 Co\textsuperscript{2+} histochemistry on the HaCaT cell line expressing TRPV1 ectopically. Cells were incubated for 10 min in buffer A containing a 20 μM CAPS+5 mM Co\textsuperscript{2+}; b 5 mM Co\textsuperscript{2+} without CAPS; c 20 μM CAPS+5 mM Co\textsuperscript{2+}+300 nM CapZ; d 20 μM CAPS+5 mM Co\textsuperscript{2+}+5 μM CapZ; e 20 μM CAPS+5 mM Co\textsuperscript{2+}+100 μM CapZ; f 20 μM CAPS+5 mM Co\textsuperscript{2+}+500 nM RuRed; g 20 μM CAPS+5 mM Co\textsuperscript{2+}+7 μM RuRed; h 20 μM CAPS+5 mM Co\textsuperscript{2+}+100 μM RuRed. The dark precipitates indicate the presence of intracellular CoS that is blockable with RuRed, a channel blocker of heat and vanilloid pain signalling. Co-application of CapZ, a competitive antagonist of pungent vanilloids, also inhibited the cellular entry of Co\textsuperscript{2+} in a dose-dependent manner; this is a well-characterized evidence-based method of localization of intracellular Co\textsuperscript{2+}. i: Gray values of the HaCaT cell line expressing TRPV1 measured by means of ImageJ software.
RuRed on TRPV1 mutants were therefore analysed with 4 μM CAPS. The neutralization of D646 reduced the sensitivity of TRPV1 to RuRed inhibition [11]. Our 45Ca2+-influx studies on TRPV1 point mutants revealed the following IC50 data for RuRed: IC50 (D646N) = 12.8 μM > IC50 (N628W) = 1.33 μM > IC50 (E651W) = 0.94 μM > IC50 (wild type) = 0.87 μM > IC50 (Y627W) = 0.26 μM (Fig. 6c), which findings correspond with published results [17]. Likewise, Co2+ inhibited the 4 μM CAPS-evoked 45Ca2+ influx in the D646N and E651W point mutants similarly as determined in the wild type. The neutralization of D646 caused no or only a minimal change in IC50. As compared with the wild type in this representative experiment, Co2+ sensitivity was slightly reduced in the D646N mutant (IC50 = 18.3 vs. 25.7 μM). The Y627W, N628W and E651W mutants displayed little or no difference relative to the wild type (IC50 = 18.6, 16.4 and 15.3 μM, respectively vs. 18.3 μM; Fig. 6d). Based on these findings, inhibition seems to be a consequence of competition with Ca2+ for M2+-chelating sites in the pore loop domain.

Discussion

Testing the effects of various metal cations on the vanilloid-induced activity of the TRPV1 channel, we demonstrated that Mg2+, Mn2+ or La3+ caused little or no decrease in
channel activity, whereas Zn$^{2+}$ proved to be a weak and only partial inhibitor of the 2 $\mu$M CAPS-induced $^{45}$Ca$^{2+}$ uptake. The other cations effectively blocked the vanilloid-induced $^{45}$Ca$^{2+}$ entry into TRPV1/HaCaT cells, with the following sequence of potency: Co$^{2+}$>Cd$^{2+}$>Ni$^{2+}$>Cu$^{2+}$.

It was reported by Nilius et al. [29] that Co$^{2+}$ reduced the inward Ca$^{2+}$ current through ECaC1 (TRPV5), a close relative of TRPV1, sharing around 30 % homology with it. Fast and reversible recovery of the current upon washout of the inhibitor was detected during their experiments. Furthermore, they identified other M$^{2+}$-s as effective inhibitors of the Ca$^{2+}$ influx. Their results indicated the following overall blocking sequence: Pb$^{2+}$=Cu$^{2+}$=Gd$^{3+}$>Cd$^{2+}$>Zn$^{2+}$>La$^{3+}$>Co$^{2+}$>Fe$^{2+}$>>Fe$^{3+}$. Zeng et al. found Cu$^{2+}$ to be a potent inhibitor of the whole-cell current evoked by intracellular ADP-ribose through TRPM2, another member of the TRP group. The inhibitory effect of Cu$^{2+}$ was irreversible, and occurred only if Cu$^{2+}$ was administered in outside-out patches, suggesting that the action site is located extracellularly. The TRPM2 current was also blocked by Hg$^{2+}$, Pb$^{2+}$, Fe$^{2+}$ and Se$^{2+}$ [30].

In accord with the above-mentioned findings, we also observed ion influx-inhibitory effects of M$^{2+}$-s on TRP channel. Depending on the TRP channel type, differences of the orders of blocking potency could be detected. Furthermore, the blocking effects of the individual cations could be reversible or irreversible, depending on the channel type. The three channels are close relatives and share high degree of sequence and structural homology with one another, which explains the similar responses to M$^{2+}$-s. Having diverged from a common ancestor, TRP channels operate on uniform principles. However, during evolution TRP superfamily has evolved for various specialized functions. TRPV1 and TRPV5, for example, belong to different subgroups of the TRP family; TRPV1–4 are non-ion-selective, whilst TRPV5–6 are highly Ca$^{2+}$-selective. Hence, this functional adaptation may cause the differences in the M$^{2+}$-evoked responses.

As concerns our own results, Co$^{2+}$ reduced not only CAPS-induced but also heat-induced $^{45}$Ca$^{2+}$ influx. When increasing concentrations of both Co$^{2+}$ and CAPS were applied, the Co$^{2+}$ inhibition patterns indicated channel-blocking kinetics. Our dose–response and Co$^{2+}$ accumulation experiments revealed a competition for binding sites and a co-entry mechanism. We presume that Co$^{2+}$ inhibits TRPV1 through its ability to bind to the ion selectivity filter of the channel: it passes through the ion channel much more slowly than Ca$^{2+}$. Ca$^{2+}$ entry is also slowed down by the binding of Co$^{2+}$, which occupies the appropriate amino acid residues of the ion selectivity filter. This hypothesis seems to be further supported by the findings of Sajadi [31], who determined the stability constants of the 1:1 complexes formed between M$^{2+}$ and L-tryptophan and other amino acids. The sequence obtained in the case of tryptophan was Ca$^{2+}$<Mg$^{2+}$<Mn$^{2+}$<Co$^{2+}$<Ni$^{2+}$<Cu$^{2+}$>Zn$^{2+}$, which follows the Irving-Williams sequence [32]. The order of the stability constants was similar in the cases of methionine, alanine, leucine, valine and glycine. The amino acid sequence of the putative pore region is STSHRWRGPACRPPDSSYNSLYSTCLELFKFTIGMGD (Q8NER1, UniProt), containing all the tested amino acids but valine. The stability constants formed between M$^{2+}$ and tryptophan were Co$^{2+}$, 2.55±0.08; Mg$^{2+}$, 2.84±0.08; Mn$^{2+}$, 3.34±0.05; Co$^{2+}$, 4.34±0.07; Ni$^{2+}$, 5.31±0.06; Cu$^{2+}$,
8.05±0.05 and Zn²⁺, 5.00±0.08. The stability constants for Mg²⁺ and Mn²⁺ are close to that of Ca²⁺, so these ions can probably readily pass through the open channel of TRPV1. Co²⁺, Ni²⁺ and Cu²⁺ can be characterized by much higher complex-forming strength, elucidating the elevated TRPV1 blocking potency. Interestingly, for these three M²⁺-s, an unexpected relationship can be observed between the stability constant and the TRPV1 blocking potency. The stronger the bond, the weaker the TRPV1 inhibition potency is. The ionic radii (in picometres) of these cations are Ca²⁺, 100; Mg²⁺, 72; Mn²⁺, 67; Co²⁺, 65; Ni²⁺, 69; Cu²⁺, 73 and Zn²⁺, 74 [33]. Ca²⁺ is likely to have the ideal ionic radius and stability constant in its reactions with amino acids in order to be effectively passed along the carbonyl groups of the peptide backbone in the ion selectivity filter and the pore loop. Co²⁺ has a medium stability constant and the smallest ionic radius, which is probably not adequate for efficient transport. These two parameters seem to be equally involved in the proper ion influx. The stronger the M²⁺-amino acid complex and the smaller the ionic radius is, the more probable it is that M²⁺ will block the Ca²⁺ influx through the TRPV1 channel. As another interesting finding, in our experiments, Zn²⁺ seemed to be only weak and partial inhibitor of the ion current. Its stability constant is almost as high as that of Co²⁺, suggesting a strong TRPV1-blocking ability, whereas its ionic radius is much larger than that of Co²⁺. Interestingly, the IC₅₀ of Zn²⁺ is the second lowest exceeding only that of Co²⁺, but Zn²⁺ can achieve a decrease of merely 30 % of the maximal ion influx.

The TRPV1 channel is a non-selective cation channel, and still shows preference for Ca²⁺. The sequence of permeability is Ca²⁺>Mg²⁺>Na⁺=K⁺=Cs⁺ [3]. In addition to all these, TRPV1 also conducts protons [34]. Following prolonged exposure to agonists, TRPV1 becomes permeable even to larger organic cations, including dyes such as YO-PRO1 and FM1-43 [22] and a lidocaine derivative QX-314 [23]. Increasing agonist concentration enhances the blocking ability of Co²⁺, suggesting a correlation between the increased efficacy of inhibition and the different open-state conformations of the TRPV1 channel. Further experiments (involving patch-clamp recordings) would be needed to clarify the inhibitory effect of Co²⁺ on TRPV1.

Positively charged tricyclics, K/R-rich basic peptides and RuRed dock to the DXExxxExXd motif at the channel orifice [13]. Although our point mutants overlap with the RuRed docking site [11, 14, 45], our point mutant studies suggest that Co²⁺ has a different binding site. Accordingly, the D646N point mutant, which is crucial for RuRed binding, does not change the kinetic parameters of Co²⁺ inhibition in cells ectopically expressing the D646N mutant TRPV1. We carried out vanilloid-induced [⁴⁵]Ca²⁺-uptake experiments with channel point mutants of TRPV1 in which the agonist binding site remained intact. Interestingly, no significantly decreased efficacy of Co²⁺ inhibition was found when 646 aspartate was replaced by asparagine. Based on these results, Co²⁺ is supposed to evoke its effect at a different site on the pore loop region than RuRed, or to use the same negatively charged amino acids passing through the TRPV1 channel as Ca²⁺. This line of research on functional point mutants should be continued to determine whether Co²⁺ has a specific binding site on the channel orifice.
Most painkiller drugs are competitive agonists and target the CAPS-binding domain [48]. As the 646DELEHTENYD acidic tetrad sequence of the TRPV1 receptor is unique among Ca^{2+}-binding proteins, this permits the design of painkillers targeting the channel orifice of TRPV1 and acting as channel blockers. A better understanding of the structural background and dynamics of the competition of Ca^{2+} with other M^{2+} for entry may result in the discovery of novel channel blocker painkillers. Furthermore, our data can contribute to a better understanding of the structures and functions of all TRP superfamily members. The specific effect of the selected M^{2+}-s on the given ion channel pore region can serve as a valuable constraint during in silico modelling of the pore region. By comparing the different cation action profiles of pore regions, the model can be fine-tuned. The mechanism of Co^{2+}-mediated inhibition provides screening for adjuvant therapeutics with higher selectivity than that of AMI, an approved drug currently used in clinical practice, but with only limited efficacy and with serious side effects.

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