TRP ion channels: Proteins with conformational flexibility

Ana Elena López-Romero, Iléana Hernández-Araiza, Francisco Torres-Quiroz, Luis B. Tovar-Y-Romo, León D. Islas, and Tamara Rosenbaum

*Departamento de Neurociencia Cognitiva, División Neurociencias, Instituto de Fisiología Celular, Universidad Nacional Autónoma de México, Mexico, Mexico; †Departamento de Bioquímica y Biología Estructural, División Investigación Básica, Instituto de Fisiología Celular, Universidad Nacional Autónoma de México, Mexico City, Mexico; ‡Departamento de Neuropatología Molecular, División Neurociencias, Instituto de Fisiología Celular, Universidad Nacional Autónoma de México, Mexico City, Mexico; †Departamento de Fisiología, Facultad de Medicina, Universidad Nacional Autónoma de México, Mexico City, Mexico

ABSTRACT

Ion channels display conformational changes in response to binding of their agonists and antagonists. The study of the relationships between the structure and the function of these proteins has witnessed considerable advances in the last two decades using a combination of techniques, which include electrophysiology, optical approaches (i.e. patch clamp fluorometry, incorporation of non-canonic amino acids, etc.), molecular biology (mutations in different regions of ion channels to determine their role in function) and those that have permitted the resolution of their structures in detail (X-ray crystallography and cryo-electron microscopy). The possibility of making correlations among structural components and functional traits in ion channels has allowed for more refined conclusions on how these proteins work at the molecular level. With the cloning and description of the family of Transient Receptor Potential (TRP) channels, our understanding of several sensory-related processes has also greatly moved forward. The response of these proteins to several agonists, their regulation by signaling pathways as well as by protein-protein and lipid-protein interactions and, in some cases, their biophysical characteristics have been studied thoroughly and, recently, with the resolution of their structures, the field has experienced a new boom. This review article focuses on the conformational changes in the pores, concentrating on some members of the TRP family of ion channels (TRPV and TRPA subfamilies) that result in changes in their single-channel conductances, a phenomenon that may lead to fine-tuning the electrical response to a given agonist in a cell.

Introduction

The appearance of the plasma membranes of cells constituted a key event for life during evolution. These structures represent boundaries of the units of life, achieving pivotal functions for the cell, including transport of molecules, communication with the environment, and metabolic functions. The intrinsic hydrophobic nature of these structures renders them impermeable to the flux of charged molecules, a process necessary for cell-cell communication, activation of signaling pathways, as well as for the ability to respond to endogenous and exogenous signals. This issue is resolved by the presence of certain types of proteins in plasma membranes, called ion channels that allow for the passage of ions from one side to the other. The specific structure of one ion channel may differ substantially from another; however, all these proteins contain a pore that opens and closes to permit the flow of ions, mediating ionic currents that, in turn, result in the generation of signals with distinct physiological consequences.

There is an intimate relationship between the structures and the functions of ion channels, with their different component regions serving a specific role in the activity of these proteins. In the 1980s, when the patch-clamp recording technique was developed, it became possible to study the behavior of single ion channels in real time[1]. Furthermore, with the advances seen in the areas of molecular biology, cloning and generation of knockout animal models, we have been provided with valuable methods[1] to carefully determine the roles of ion channels in normal and under pathophysiological conditions. A large variety of ion channels have been cloned and biophysically characterized and, in the past three decades,
research-oriented towards determining the structure of ion channels has witnessed remarkable progress.

The Nobel Prize in Chemistry was awarded to Roderick MacKinnon and Peter Agre in 2003, for advancing the field by resolving the molecular structures K⁺ channels [2–5] and aquaporins [6,7], allowing us to further understand their functional features.

The present review focuses on a family of proteins named Transient Receptor Potential (TRP) ion channels, which has fundamentally reshaped our knowledge of sensory physiology and their importance is illustrated by their pivotal roles in smell, taste, vision, touch and our ability to detect changes in temperature. Originally, the first trp gene was identified in a Drosophila mutant with altered vision [8]; however, it was not until it was cloned that the deduced amino acid sequence led to the suggestion that trp encoded for a cation channel [9]. Electrophysiological studies later showed that trp indeed was an ion channel and that its selectivity could be modified by inserting mutations in the amino acid region that formed the pore loop [10].

The field of study of TRP channels witnessed a boom when mammalian homologs of the Drosophila trp channel were cloned and when some of these were identified as temperature sensors [11,12]. A feature of these proteins is that several of these channels are polymodal, that is, being activated by several distinct physical stimuli and more than one ligand. In some cases, different biophysical properties as well as distinct conformational changes in their pores, associated with interactions with different ligands, have been demonstrated.

We will discuss a few examples of ion channels where conformational changes are associated with different ion conductance states. However, we will mainly focus on TRPV1 and other TRP ion channels for which structures have been solved, making emphasis on the conformational changes produced by various ligands and on their effects in the functional properties of these proteins.

**Rearrangements in the outer pore of kv2.1 lead to changes in ion conductance**

For voltage-gated ion channels, modulation of macroscopic current magnitude archetypically is thought to occur through an effect on the gating (opening and closing properties) of these proteins. In this sense, most studies of gating have concentrated on the voltage-dependent gate at the cytoplasmic entrance to the pore. Moreover, there is a large body of evidence that also supports an important role for the selectivity filter in gating activation [13,14]. It has been suggested that some form of voltage-dependent gating can exist at the selectivity filter (the region that discriminates among types of ions passing through the pore) [15–17], although the main closed-open transition is controlled by the S6 bundle-crossing intracellular gate [18].

These aspects have been explored in Kv2.1 potassium channels, which are slowly inactivating delayed rectifiers present in non-neuronal excitable cells and several neurons. In these channels, the current magnitude is modulated by the external K⁺ concentration, making outward currents through Kv2.1 channels become larger when the extracellular K⁺ concentration is increased [19]. Kv2.1 channels exhibit two distinct pharmacological profiles as a function of the K⁺ concentration since they can either be sensitive to external tetraethylammonium (TEA; IC₅₀ ~ 3–5 mM), or completely insensitive to this blocker [20]. The underlying mechanism of these effects encompasses the opening of Kv2.1 channels into one of two different outer vestibule conformations with different sensitivities to TEA. It has been shown that the channels that open into a TEA-sensitive conformation produce larger macroscopic currents [19]. In contrast, channels that open into a TEA-insensitive conformation, a phenomenon that occurs in the presence of higher potassium concentrations, yield smaller macroscopic currents [19].

Trapani and collaborators examined the mechanism by which conformational changes in Kv2.1 channels produced changes in current magnitude. By using a combination of single-channel, macroscopic recordings, and hidden Markov modeling they proposed a model in which an outer vestibule lysine residue in position 356 (pore turret region) interferes with the flux of K⁺ through the channel [19]. This led to the suggestion that the K⁺ concentration-dependent change in orientation of that specific K356 can modify single channel conductance through a change in the level of such an interference [19].
In summary, in the conformation where currents are smaller in magnitude, K356 would be oriented towards the center of the ion conduction pathway, not easily allowing the flux of K$^+$ ions and resulting in a change in the single-channel conductance. In contrast, when K356 is oriented away from the conduction pathway, it readily allows the passage of ions. Thus, these results obtained by Trapani et al., not only provided evidence where Kv2.1 single-channel conductance is modulated by the outer end of the conduction pathway through the occupancy of open states with different outer vestibule conformations, but they also showed that these occur under physiologically-relevant K$^+$ concentrations [19].

Hence, in a physiological scenario, it is important to consider that with rising extracellular K$^+$ concentrations, the current magnitude of K$^+$ channels in most neuronal cells is reduced as a result of a decrease in electrochemical driving force. However, Kv2.1 counteracts this reduction in net outward current in the presence of higher K$^+$ concentrations and, as suggested by the authors, this would provide the cells where these channels are expressed with a mechanism that maintains action potential integrity when high-frequency firing conditions are present [19].

The validity of the pore dilation phenomenon to explain changes in ion channel conductance

Ion channels exhibit different permeability and selectivity to various ions. Potassium, sodium, and calcium channels contain ion binding sites in their selectivity filters that enable them to finely discriminate among the ions that will flow through their pores[21]. Nonetheless, some ion channels have been shown to exhibit dynamic changes in their ion selectivity in response to agonist activation that could, in theory, allow for changes in their conductance. Examples of these ion channels are P2X receptor channels [22,23], acid-sensing ion channels (ASIC) [24,25] and TRPV1 channels [26,27]. This phenomenon has been called pore dilation and it will be next discussed for P2X receptors, a well-studied example of this process, and for TRPV1 in a later section.

P2X receptors are ion channels gated by extracellular ATP [28] and it was originally thought that the pore first opened rapidly to a conducting state selective for small cations such as Na$^+$, K$^+$ and Ca$^{2+}$ (I$_1$ current) and gradually increased in size or dilated over time, rendering the channel permeable to large organic cations such as N-methyl-D-glucamine (NMDG$^+$) or I$_2$ current, as well as to large fluorophores and dyes (i.e. YO-PRO1) [22,23]. However, based on the original assumption that P2X receptors experience a slow, time- and agonist-dependent pore dilation, three mechanisms have been suggested to explain this phenomenon: (1) it represents an intrinsic gating property of the functional P2XR channel, where ATP exposure results in the widening of the pore and a change from a state that conducts Na$^+$, K$^+$, Ca$^{2+}$ (I$_1$) to one that conducts NMDG$^+$ and YO-PRO1, a large propidium dye. Such a switch has been proposed to be modulated by phosphorylation or dephosphorylation events [22,23]; (2) macropores are formed as a result of an agonist-dependent redistribution and oligomerization of P2XRs. The fusion of two or more trimeric P2XRs, as well as an enlargement of the channel pore, or the formation of a separate but larger pore among aggregating trimeric assemblies, could underly macropore formation. This hypothesis has been ruled out by some research groups since no clustering or redistribution of channels expected during oligomerization has been found to occur [23,29], but some others still support it [30]; (3) the permeability to large cations is facilitated by a structurally separate transport pathway downstream of P2XR activation [22].

Although the phenomenon of pore dilation has been extensively explored, accumulating evidence suggests that it may be an artifact of the experimental conditions used, as will be detailed in the Discussion section.

Next, we will discuss some structural characteristics of TRP ion channels that shed light into why these proteins represent an example of conformational flexibility in response to agonists that may result in changes in cellular excitability.

The structures of thermotgrp channels reveal conformational flexibility

The TRPV1 channel was the first mammalian TRP channel to be cloned. In 1997, it was shown that
TRPV1 is the receptor for capsaicin [11,12]. In this groundbreaking study, the laboratory of David Julius isolated a cDNA clone that reconstituted the response to capsaicin in non-neuronal cells. By examining the amino acid sequence of this clone, they demonstrated TRPV1 is an integral membrane protein, homologous to a family of putative store-operated calcium channels and that it is expressed by small-diameter neurons within sensory ganglia such as the dorsal root (DRG) and trigeminal (TG). Moreover, these authors showed that TRPV1 is also a thermal sensor, activated in response to temperatures in a range known to elicit pain-associated behaviors in animals and pain in humans [11]. It was later shown, in 1998, that TRPV1 is also activated by low extracellular pH (pH ≤ 5.9) [12].

Continued work on the study of the function of TRPV1, one of the best studied TRP channels, has led to the identification of several other agonists and modulators including resiniferatoxin (RTx) [12], diacylglycerol [31], hydroperoxyeicosatetraenoic acid (HPETE) [32], anandamide [33], N-acyl ethanolamines (NAEs) [34], n-acyl-dopamines [35], nitric oxide [36], cations [37,38], hydrogen sulphide (H2S) [39], the double-knot toxin (DkTx) from the Earth Tiger tarantula [40] and lysophosphatidic acid (LPA) [41].

TRP channels are all tetrameric structures. The structure of TRPV1 had been initially solved in the year of 2008 by the group of Vera Moiseenkov-Bell. These authors obtained a 19 Å resolution cryo-EM structure that showed that TRPV1 exhibits a four-fold symmetry with a large open basket-like domain, formed by the N- and C-termini of the channel as well as a compact transmembrane region [42]. In this first solved structure of TRPV1, the authors described a 150 Å tall ion channel which is divided into a “small” compact region (30% of the total volume) and into a “large” region or the “basket-like” domain (70% of the total mass).

This seminal study was followed by the higher resolution structures for TRPV1 obtained by the research groups of Yifan Cheng and David Julius. Two, back-to-back reports, described structures of 3.4–4.2 Å resolution obtained by cryo-EM [43,44]. They confirmed that TRPV1 has a four-fold symmetry with a central ion pathway that is formed by transmembrane segments S5 and S6 and a pore-loop, all of which are surrounded by the S1-S4 domains with a domain-swapped geometry [43,44]. The pore region was described as one containing a wide extracellular mouth and a short selectivity filter. The important TRP domain, which is a region conserved in these family of ion channels and that is thought to play an important role in their allosteric modulation, was shown to interact with the S4-S5 intracellular linker. Each of four subunits in TRPV1 contains six ankyrin repeats that form the ankyrin repeat domain (ARD) localized to their N-termini [43,44] (Figure 1).

These research groups also described the structures of TRPV1 obtained in the presence of different agonists, providing insight into the mechanisms that allow the polymodal activation of this channel. One of these structures was obtained in the presence of RTx (which also binds to the vanilloid pocket where capsaicin binds through residues Y511 and S512 in S3 and M547 and T550 in S4) and DkTx together. The authors showed that DkTx was bound to the extracellular loops of the outer pore region and defined contacts of the toxin with residues at the top of the pore helix of one subunit and the outer pore loop in the proximal S6 region of another nearby subunit [43,45] (Figure 2).

The structures of TRPV1 obtained in the presence of both agonists display evident rearrangements in the ion conduction pathway and the outer pore region. With RTx and DkTx bound to the channel, the ion conduction pathway appears completely eased of constrictions. This is in contrast to the structure obtained only in the presence of capsaicin, where the lower gate I679 is extended, but the selectivity filter G643 remains unaltered [45] (Figure 2).

When TRPV1 has RTx and DkTx bound, a shift or tilt of the S6 helix 1.9 Å away from the central axis of the channel is produced, as compared to the apo or unliganded state. Moreover, an increase in the distance between the carbonyl oxygens of G643, which constitutes the narrowest point of the selectivity filter, is also observed when both ligands are present: it goes from 4.6 Å in apo (PDB 3J5P) to 7.6 Å in RTx/DkTx (PDB 3J5Q) [45].

As mentioned above, G643 (at the upper gate) remains unaltered between the apo structure (PDB 3J5P) and in the structure in complex with
capsaicin, and this could be explained because, in the absence of DkTx, the channel could be undergoing transitions to closed states. However, the distance between the I679 residues (lower gate) in the apo state is 5.3 Å (PDB 3J5P) and, when in complex with RTx/DkTx (PDB 3J5Q), it shifts to 9.3 Å, while it is at 7.6 Å with capsaicin only (PDB 3J5R). Therefore, the authors concluded that the structure obtained with capsaicin is likely only a partially activated state of TRPV1 [45].

When TRPV1 is in the presence of capsaicin only, the structure can be superimposed to that of the apo state in the outer pore region. Hydrogen bond interactions among side chains of the E600 residues and main-chain nitrogen atoms of Y653 and D654 are broken in the structure with RTk/DkTx but maintained with capsaicin only.
that stabilize the closed state change when the channel is in the apo state or in the presence of RTx/DkTx: D654-E600 goes from 3.6 Å to 8.4 Å, while for Y653-E600 it goes from 4.4Å to 8.3Å, respectively [45] (Figure 2).

In summary, this ion channel displays extraordinary flexibility of movement in response to the presence of different agonists.

Another example of conformational changes that may allow for “superactivated” states is that of TRPV2. This channel is activated by temperatures near 52ºC [46] and by membrane stretch [47]. Although structurally similar to TRPV1, TRPV2 is not activated by vanilloid compounds. However, the laboratory of Kenton Swartz produced a TRPV2 channel with four-point mutations to form a vanilloid-binding site located in the S4-S5 linker and the base of S5 [48], rendering this channel sensitive to RTx. These results were also confirmed by Yang et al., who used a TRPV2-Quad channel (with the same point mutations in mouse), where capsaicin competed for the same site as RTx [49]. This TRPV2_Quad channel was found to bind RTx, and that leads the channel to an unstable open state through a mechanism that bridges the S4-S5 linker to the S1-domain [49].

A couple of years later, Zubcevic and collaborators obtained the crystal structure of a minimal construct of the rabbit TRPV2 channel (miTRPV2) in the absence of agonist and compared it with that of the vanilloid-sensitive miTRPV2 (F470S, L505M, L508T, and Q528E, in the rabbit TRPV2) in complex with RTx [50].

Interestingly, in the structure in complex with RTx, they found that two subunits adopt distinct orientations between the S4-S5 linker and the S5, as compared to the other two subunits. In other words, while two subunits (A and C) are found to spread away from each other, the other two (B and D) get closer. This rearrangement results in an asymmetric pore that adopts a wider conformation at the selectivity filter when compared to the structure without RTx (Figure 3).

The authors determined that with RTx the distance between the G604 carbonyl oxygens is ~7.1 Å, and in the widened subunits the distance between the I603 carboxyls was ~12.3 Å, while in the structure with only Ca$^{2+}$ the distances between G604 carbonyl oxygens are 6.1–6.4 Å [50,51]. The conclusion was that, when RTx binds to the channel, it pushes the base of S5 downwards and produces a bend in the S4-S5 linker producing a different conformation that leads to rotation and widening of the entire involved subunit (Figure 3).

The authors also suggested that this wider conformation of the pore could allow for the permeation of larger cations since they performed experiments where it is observed that, upon activation of the vanilloid-sensitive TRPV2 with RTx (250 nM), uptake of the large fluorescent molecule YO-PRO-1 (376 Da) could be attained. With this experiment, they confirmed the functionality of this conformation, and suggested that it was either a fully open state or an intermediate state that occurred before a symmetric open state capable of permeating large organic cations [50].

Just as in TRPV2, a vanilloid-binding site can be introduced into the TRPV3 channel upon the insertion of six residues in the putative vanilloid-pocket: W521Y, H523E, F522S, L557M, A560T, and Q580E, as shown for the mouse TRPV3 channel by Zhang and collaborators [52]. Nonetheless, the insertion of this vanilloid-binding site is not enough to achieve activation of TRPV3 by RTx and the presence of other agonists, such as 2-APB (aminoethoxydiphenyl borate) or heat, is required [52]. These results demonstrate that gating pathways are conserved among the TRPV channels, but they also show that the energetics for activation can dictate their sensitivity to certain stimuli since adding a binding site for an agonist does not necessarily result in the activation of the ion channel [52].

Structurally, the sensitized state of the human TRPV3 undergoes an α-helix to π-helix transition in the S6 which, in turn, disrupts the S4-S5 linker and S6 and the latter bends ~9° away from the pore [53]. In a previous report, the open conformation of the mouse TRPV3 obtained with 2-APB only shows a π-helix in the S6 [54], leading the authors to suggest that the secondary structure transition could be a hallmark for a sensitized channel, in other words, a state that requires less energy to activate [53].

Additionally, Zubcevic and collaborators also observed that during the activation of TRPV3
with 2-APB, the intermediate states display asymmetrical rearrangements that are not present in the apo or sensitized states. The differently observed deviations for the subunits are caused because the \( \pi \)-helices in the S4-S5 linkers begin at different residues. This is interesting because it accounts for a similar phenomenon as the one described above for the activation of TRPV2 with RTx, which also originates at the S4-S5 linker [50]. Together, these results further support the hypothesis that the gating mechanisms of TRPV channels are conserved and a break in symmetry may be characteristic to these channel’s gating.

The TRPV4 channel was first described as an osmosensor, capable of activating after exposure to extracellular hypotonic conditions [55,56]. Subsequent reports demonstrated that, like other members of the TRPV subfamily, TRPV4 is a heat-sensitive channel activated by temperatures above 27°C [57] and chemical agonists such as phorbol derivates [58] and 5,6-epoxyeicosatrienoic acid, a metabolite of arachidonic acid [59,60].

Deng and collaborators obtained the first high-resolution cryo-EM structure (3.8 Å) of the TRPV4 channel [61]. These authors determined that the channel was in a closed conformation, that the narrowest diameter in the pore (5.3 Å) was located at residue M714, which impeded the permeation of ions. Even in the closed state, TRPV4 displayed a particularly wide selectivity filter: the diameter at G675 in TRPV4 was 10.6 Å, while the diameter at G643 in TRPV1 in complex with RTx/DkTx was 7.6 Å [45]. This evidence suggested that unlike TRPV1, the TRPV4 channel lacks an upper gate [61].

The single-channel conductance of TRPV4, when stimulated with hypotonic solutions, has been reported to be around 310 pS, at positive voltages [55]. However, reports for its single-channel conductance during spontaneous activity is near 88 pS [56], which is near the value (around 98 pS) obtained with other agonists such as 4α-Phorbol 12,13-didecanoate (4αPDD) and with heat (around 105 pS) [58]. Altogether these results

Figure 3. Comparison of open and closed states of TRPV2.
Extracellular views of A) a closed miTRPV2 channel (PDB 6BWM) and B) vanilloid-sensitive miTRPV2 (PDB 6BWJ) in complex with RTx shown as yellow spheres, representing the open state. On the lower panel are side views of the closed (C) and open (D) channel. Only two subunits are shown for simplicity. The green sphere is Ca\(^{2+}\) trapped in the pore which is notably wider in the open configuration, its also visible the rearrangement of the intracellular domain. The side chains of G604 are also depicted.
suggest that the channel can adopt a different functional state when it is stimulated under hypotonic conditions vs. heat or phorbol, and that differences in the conductance of TRPV4 may be observed with other agonists, but further studies must be performed.

TRPA1 is a channel mainly expressed in nociceptive neurons which responds to multiple noxious stimuli, including pungent chemicals present in garlic, mustard oil, cinnamon or wasabi [62,63]. Moreover, TRPA1 was initially described as a channel activated by noxious cold temperatures [64,65], but there has been controversy around this topic since this channel functions as a temperature receptor in a species-specific fashion [66–68]. Chemical agonists of TRPA1 have been divided based on their mechanism of action into electrophilic and non-electrophilic agonists. Examples of the first are allyl isothiocyanate (AITC) [69] or allicin [62] that activate the channel through covalent modification of cysteines at the N-terminal domain. In fact, allicin can also activate TRPV1 through the same mechanism [70–72]. The non-electrophilic agonists include carvacrol, oleananthal, Δ9-tetrahydrocannabinol (THC) and acidic pH, which activate TRPA1 by mechanisms not associated with cysteine modifications and which remain mostly unclarified [73–75].

Cavanaugh et al., determined, in excised TRPA1-expressing membrane patches, that the channel needs to be exposed to polyphosphates (PPPi) [76] in order to be activated by AITC, but in this AITC-insensitive conformation (in the absence of PPPi) it can be activated by THC [77]. This result indicates that the channel shifts to a different functional conformation unresponsive to cysteine modification but responsive to activation by a different mechanism [77], each with a distinguishable function and agonist dependence.

The different functional states described by Cavanaugh et al., have not been structurally compared. Unlike TRPV1, the cryo-EM structure for TRPA1 was not obtained in the presence of different agonists, but only with AITC (PDB 3J9P). Nonetheless, it was possible to determine that the pre-S1 helix, linker S4-S5, and TRP-like domain are bound by hydrophobic interactions and represent an important site for allosteric modulation, similar to what happens with TRPV1 [45,78].

Nevertheless, recent studies have compared the structural rearrangements of TRPA1 in the presence of different agonists using techniques such as limited proteolysis combined with mass spectrometry [75] or circular dichroism spectroscopy only for the N- and C-terminal domains [79]. These robust methodologies show important interactions between cytoplasmatic domains involved in the gating of the channel [79], as well as differences between the activation with electrophilic and non-electrophilic agonists [75].

TRPV5 and TRPV6 display limited structural rearrangements in their pores

Both TRPV5 and TRPV6 ion channels are expressed mainly in the epithelial tissue of the digestive tract and kidney, where they play roles in the reabsorption and transcellular transport of Ca²⁺ [80]. Unlike other members of the TRPV family, TRPV6 and TRPV5 show inward rectification and these highly Ca²⁺-selective proteins are constitutively active under physiological conditions [81]. Despite the high ion selectivity, in the absence of extracellular divalent cations, they can permeate monovalent cations [80].

TRPV5 and TRPV6 channels share structural traits that differentiate them from the other known TRPV channels. Three phenylalanine residues in S6 give rigidity to the pore, and a ring of aspartates in the selectivity filter confers its high Ca²⁺ selectivity [82]. Neutralization mutation of the negative aspartate charge not only affects Ca²⁺ selectivity but reduces inward rectification [80].

Functionally, both channels can be inhibited by endogenous calmodulin (CaM) [83] or by econazole, an antifungal [84], while PIP₂ helps stabilize their open state [85]. The conformational changes between open and closed states of these channels have been explored by comparing the cryo-EM structures of econazole-bound TRPV5 and of both channels in the presence of either PIP₂ or CaM [86,87].

One of the main findings of this comparison is that the selectivity filter remains mostly the same regardless of the molecule that binds the channel. The lower gate identified in TRPV5 is formed by F574, M578, and H582 that constrict the pore to a diameter between 4.5 and 5.9 Å in the same bundle crossing configuration observed in other TRP channels. The closing seems to involve a change in the position of the S4-S5 linker and the loop connecting S6 with the TRP domain [86,87].
The binding of PIP$_2$ induce a change in the orientation of the lateral chain of the aspartate in the selectivity filter. This change may be explained by an interaction of the head group of PIP$_2$ with R584 in the S6 helix, analogous to what has been observed in TRPV1, which rotates and pulls away from the center of the pore accompanied by movement of the S4–S5 linker [87]. The main difference in the response of TRPV5 and TRPV6 was observed in the CaM-bound structures. Both channels bind to a single CaM-molecule which is capable of physically blocking the pore from the intracellular side, but while TRPV5 shows no conformational changes, TRPV6 shows an α- to a π-helical transition of S6 which tilts the helix toward the center, so I575 becomes an inactivation gate [87], which has also been observed in TRPV3 [53].

Analysis of the pores of TRPV5 and TRPV6 points to rigid structures that contrast with the more flexible pores of TRPV1–4. To date, there is no physiological evidence of changes in their single-channel conductances in response to different stimuli, but it has been suggested that their activity might be regulated by the formation of TRPV5/V6 heterotetramers, as they are co-expressed in several tissues. Hoenderop et al. used *Xenopus laevis* oocytes to express such heterotetramers and found that they have Ca$^{2+}$-dependent inactivation and block by ruthenium red with features intermediate to those of TRPV5 and TRPV6 [88]. Although this is an interesting possibility, they have not been identified in vivo, and their biophysical properties at the level of single-channels have not been investigated.

**Lysophosphatidic acid: A pain-producing phospholipid**

Another activator of TRPV1 shown by our group to directly interact with this ion channel is lysophosphatidic acid (LPA) [41]. This is the only example for which a change in single channel conductance in a TRP channel has been shown to depend upon the presence of different agonists; thus, we will discuss this phenomenon in detail.

LPA has been extensively linked to the generation of chronic neuropathic pain through its interactions with G protein-coupled receptors (GPCRs) [89–92] and to modulate the activity of several ion channels [93–97]. Regulation of the activity of TRPV1 by molecules of a lipidic nature has been explored, and it has been described that PIP$_2$ [98–106], polyunsaturated fatty acids (PUFAs) [107], monounsaturated fatty acids [108] and cholesterol [109–111] can modulate the activity of TRPV1, either directly or indirectly. However, reports of direct interactions of LPA with ion channels, in general, are scarce [95,112,113].

LPA is a phospholipid constituted by an acyl chain of saturated or unsaturated fatty acids associated with a glycerol backbone by ester links, and that contains a phosphate head group [114]. LPA can be naturally found as different species that vary in their acyl-chain length as well as in their saturation (16:0; 18:0; 16:1, 18:1; 18:2 and 20:4).

Our group studied whether LPA could produce acute pain in mice and showed that it is considerably dependent upon TRPV1. These results led us to determine that LPA could, in fact, produce currents and action potentials in DRG neurons from wild-type mice but not in DRG neurons from *Trpv1$^{-/-}$* animals. Using a heterologous expression system of HEK293 cells transfected with TRPV1, we found that currents could be induced when LPA was applied to the intracellular or extracellular faces of the ion channels and that the responses were dose-dependent [41].

In our studies, we ruled out the participation of GPCR-related signaling pathways on TRPV1 activation by LPA and identified a site of interaction for this phospholipid, residue K710 in the C-terminus of TRPV1 [41]. Confirmation of these results was obtained later by another group in a study where TRPV1 was expressed in lipid bilayers and activated by LPA [115].

Hence, we extended the aforementioned study and determined that, for fatty acids to activate TRPV1, the following features had to be met by these molecules: the presence of a charged phosphate group, a length of, at least 18 carbons of the acyl chain and a mono unsaturation [116].

**LPA produces a different open conformational state to that of capsaicin**

Our first study on the effects of LPA on TRPV1 was published in 2011 [41], and since then, we have been studying the specifics of how this molecule regulates the activity of TRPV1. Our most recent study on this subject provides functional
evidence that LPA produces a different open conformational state of TRPV1 that has a larger conductance.

In preliminary experiments, we had found that addition of LPA (5 μM) to excised membrane patches of TRPV1-expressing HEK293 cells resulted in larger magnitude macroscopic currents, as compared to those obtained in the presence of saturating capsaicin concentrations (4 μM). We had also observed that LPA (5 μM) produced activation of TRPV1 single-channel currents with an open probability (Po is 0.8) similar to that of capsaicin (4 μM).

These data could be explained as 1) LPA either producing an increase in the number of ion channels in the excised membrane patches (which was highly improbable); 2) An increase in the membrane negative surface charge near the channel leading to an accumulation of positively-charged ions near the pore mouth of TRPV1, 3) Producing a phenomenon termed “pore-dilation” in which the permeability to large ions occurs in the presence of prolonged exposures to agonists and that had only been described for activation with capsaicin [26] and/or 4) a change in the single-channel conductance.

In this last study, we performed a set of experiments in order to discern among all of these possibilities. We started by studying how LPA affected the unitary TRPV1 currents by first applying capsaicin (4 μM), washing off the agonist and then applying LPA (5 μM) to the same membrane patch. The results showed that LPA produced a 41% increase in the magnitude of single-channel currents at a single voltage (60 mV), which was also observed at other voltages (Figure 4).

Furthermore, our experiments ruled out all of the other possibilities enlisted above and showed that coapplication of subsaturating concentrations of both agonists together resulted in two different types of openings in a single channel, with conductances corresponding to those achieved either by adding capsaicin or LPA alone [117]. This was the strongest indication of these two agonists producing distinct open conformations that varied in their conductance. Finally, we defined that the presence of K710, the residue that is pivotal for the activation of TRPV1 by LPA, was also necessary for the generation of the higher-conductance open state [117].

**TRPC and TRPM channels**

Until just recently, finely defined structural characteristics of TRPC channels have remained elusive, mainly because of the lack of high-resolution structures. The cryo-EM structures obtained to date are those for the TRPC3, TRPC4, TRPC5 and TRPC6 channels [118–123]. However, no functional studies reporting changes in the conductance of these channels in response to different agonists are available.

These channels exhibit four ankyrin repeats and linker helices at their N-terminal domains and at the C-terminal domains they possess a TRP domain that connects a helix and a coiled-coil domain. Inside the pore, all TRPC members have a conserved “LFW” motif that maintains the structure of the pore [124].

---

**Figure 4.** LPA increases single channel currents. Current amplitude for a TRPV1-expressing membrane patch in the presence of capsaicin (4 μM; left) and in the presence of LPA (5 μM; right). Traces were obtained at +60 mV.
The TRPC subfamily can be divided into two groups that share some distinctive characteristics. The first group is constituted by the TRPC3/C6/C7 channels that are activated by DAG. Structurally, both TRPC3 and TRPC6 display a pre-S1 elbow, a TRP re-entrant helix embedded in the membrane bilayer after the TRP domain, and an unusually long S3 that forms a “pseudo” extra cellular domain (ECD) [121,122,124]. Furthermore, unlike most TRP channels, TRPC3 channels possess a linker that connects the TRP domain to the S6. This results in a more flexible TRP domain that exhibits a higher coupling with the S4-S5 linker and a lipid binding site near the pre-S1. All of these peculiar interplays of domains may provide a molecular basis for TRPC3 lipid gating [121].

The second group includes the TRPC4/C5 channels that are modulated by Gq, Gi/o GPCRs, membrane lipids, and intracellular Ca2+ [119,125]. These channels share the common feature of a disulfide bond between two cysteines at the loop linking S5 and the pore domain that maintains the residue E555 in proper position and thus stabilize the upper region of their selectivity filters [119,120,123,124]. As in other TRP channels, the TRP domain of TRPC4 is an important molecular effector for gating in this protein since it interacts with the S4-S5 linker and other proteins [123].

The structures of TRPC3/C6 and TRPC4 (both human and mouse) show a closed pore conformation. For TRPC3 the lower gate is located at residues I658 and L654 where they present a radius of less than 1 Å [121]. As for the human TRPC4, the residues I617 and N621 were identified as the lower gate where the constriction of the diameter was around 1.6 and 0.7 Å (defined by opposing van der Waals surfaces) [123], and for the mouse TRPC4 the narrowest point is located at N621 (3.6 Å) and constitutes the lower gate along with I617 and Q625 [119].

However, the TRPC5 is considered at least partially open since the constriction at the lower gate formed by the residues I621, N625 and Q629 was around 4.9 Å [120]. Comparing this structure with the closed TRPC4, Duan and collaborators suggest that the extracellular disulfide bond is a transducer of conformational changes. Nonetheless, more structures of TRPC channels at their open conformation will help to elucidate if there are differences in the pore during their gating [120].

The TRPM subfamily is composed of eight members with diverse features and functions [126]. The TRPM members for which cryo-EM structures have been solved in the closed or partially open conformations are TRPM2, TRPM4, TRPM7 and TRPM8 [127–135]. Their subunits are composed of six transmembranal segments, but at their N-terminal domains, they present four TRPM homology regions (MHR) and a C-terminal coiled-coil domain [127–135].

The C-termini of TRPM2, TRPM6, and TRPM7 channels possess enzymatic domains and have been named “chanzymes” [136–139]. Similar to TRPC4/C5 channels, TRPM channels possess disulfide bonds at their extracellular pore that stabilize the pore domain, but these disulfide-bonds form between cysteines at the pore loop and the pore helix S6 [119].

TRPM2 is a non-selective cation channel sensitive to temperature [140] that is activated when ADP-Ribose (ADPR) and Ca2+ are co-applied [141]. Wang and collaborators obtained the structures of human apo-, ADPR-bound, and ADPR-Ca2+-bound TRPM2 channels. They observed that the binding of ADPR alone produced a closed conformation, considering it a “primed state” [128].

The lower gate was identified at residues I1045 and Q1053 in the pore and shown to be enlarged in the ADPR/Ca2+ structure. Although a change distance of the residues that form the gate was observed, it was not considered large enough to allow passage of Ca2+ (~4 Å radius) [128].

For zebrafish TRPM2 channels it was shown that, once ADPR binds to the MHR1/2 domain, a displacement of MHR4 is transmitted to the TRP helix, pushing it towards the S4-S5 domain. Consequently, the S4-S5 relocates the S5 and S6 and finally produces the opening of the channel. Again, as for other TRP channels, the TRP helix is shown to act as a transducer between the intracellular and transmembranal domains during gating [129]. Moreover, the authors propose putative binding sites for Ca2+ in the S3, leading them to speculate that Ca2+ facilitates the opening due to the movement of the S3 upon the binding of this ion which allows for the relocation of the S4-S5 linker [129].

TRPM4 and TRPM5 channels are among the only TRP members that are not permeable to
divalent cations [132,142,143]. Despite their ion selectivity, TRPM4/5 channels are activated upon intracellular Ca\(^{2+}\) increases [144] and can be modulated by PI(4,5)P\(_2\) [145,146]. However, TRPM4 is inhibited by intracellular ATP and nucleotides, while TRPM5 is insensitive to these molecules [147,148].

The pore of TRPM4 has two gates: one located at the selectivity filter and another formed by the intracellular portion of the S6. In the human channel, the selectivity filter gate is formed by residues 975-FGQ-977 [130–132]. Likewise, the intracellular gate is formed by the I1040 residue [131,132]. Strikingly, both gates remain unchanged in all reported Ca\(^{2+}\)-bound structures, and it has been suggested that Ca\(^{2+}\)- binding is required for the opening of the channel in response to voltage [131].

In the structure obtained in complex with ATP, Guo and collaborators located a nucleotide binding domain along with twelve helices and two ankyrin repeats in the N-terminal [149]. The ATP-bound structure is an inhibited state of TRPM4, and there were no overall changes in the pore compared to the apo structure. The TRP domain forms hydrophobic interactions with the S4-S5 linker and forms a cavity between the S1-S4 domain, which harbors a potential Ca\(^{2+}\)-binding site [149].

The protein kinase domain in TRPM7 induces phosphorylation of receptor tyrosine kinase (RTK)-signaling intermediates and chromatin modifications [150,151]. TRPM7 is permeable to Mg\(^{2+}\), Zn\(^{2+}\) and Ca\(^{2+}\) [152], and the cryo-EM structure of TRPM7 was obtained in the presence and absence of Mg\(^{2+}\) [133].

There were no notable changes on the pore domain of TRPM7 under different ionic conditions. Interestingly, the selectivity filter identified in TRPM7 (1045-FGE-1047), varies only on one residue to the selectivity filter described in TRPM4 (971-FGQ-973 in mouse TRPM4 channel), which is crucial to confer the monovalent selectivity of the latter. The lower gate was determined to be formed by I1093 and N1097 residues [133].

The TRP domain of TRPM7 interacts with different regions of the channel, highlighting the importance of this region for gating in this protein. For example, the TRP domain interacts with the N-terminal residues S744 and Q740, with the S4-S5 linker by forming hydrogen bonds between W1111 and R1115 (both located to the TRP domain) and with A981 and V982 (in the S4-S5 linker) [133]. Additionally, it establishes π-π stacking (Phe1118/Tyr1122) and cation-π interactions (Arg1115/Trp1111) among its α-helical structure [133]. However, unlike what has been observed in other TRP structures, the TRP domain of TRPM7 does not contact the pre-S1 helix [133].

Another region that may produce important conformational rearrangements in TRPM7 is the pore helix. It has been shown that a disulfide-bond established between residues C1056 and C1066, is only formed in the presence of Mg\(^{2+}\). Thus, it has been proposed that this cation could stabilize the bond and impact directly on the structure of the pore [133].

Finally, the TRPM8 is a non-selective cationic channel permeable to Ca\(^{2+}\) and activated by cold temperatures (below 25°C) and menthol [153]. The channel is modulated by PI(4,5)P\(_2\) [153,154].

In the same report, the cryo-EM structures for TRPM8 have been obtained in complex with PIP\(_2\), Ca\(^{2+}\) and icilin, and with WS-12 (a menthol analog) together with PIP\(_2\) [134]. These structures demonstrate that both, icilin and WS-12, bind to a cavity formed between S1-S4 and the TRP domain [134].

The structures reveal a different binding site of PIP\(_2\), as compared to other TRP channels. This site is formed by the pre-S1 domain, the S4-S5 junction, the TRP domain and the MHR4 of an adjacent subunit. Interestingly, upon the binding of PIP\(_2\), the S4 undergoes a change from a complete α-helical to a 3\(_{10}\) helical conformation [134]. This produces the movement of the TRP domain and the S5 toward the cavity. At the same time, it disrupts the interaction among the S1-S4 domain and the S6-pore domain, enabling gating [134]. Additionally, during gating, the S1-S4 domain of TRPM8 suffers a rigid rotation away from the pore [134], unlike what happens to the S1-S4 domain in TRPV1 which remains static [45].

The lower gate of TRPM8 is formed by L973 where the tightest constriction along the pore was found in a previous structure [134]. Yin and collaborators determined that both structures (PIP\(_2\)-Ca\(^{2+}\)-icilin and WS-12-PIP\(_2\)) were non-conducting states. Although, in the icilin-PIP\(_2\)-Ca\(^{2+}\) structure the S6 is curved, which
suggested that there may be a π helical turn along the S6 as previously described for the sensitized state of TRPV3\(^5^3\). Therefore, the structures may represent sensitized or presensitized states of TRPM8 [134].

Although conformational changes in TRPM channels in response to agonists or modulators of their activities have been clarified, changes in their conductance levels in response to these molecules have not been reported.

**Discussion**

Here we have summarized some phenomena that could potentially lead to changes in the conductance of single ion channels. For example, pore dilation is a phenomenon suggested to occur in TRPV1 [26,155] and other channels such as the P2X receptors, and it is characterized by a change in the permeability of the ion channel to larger ions, induced by long times in the presence of the agonist. However, the results that we have obtained, as well as those of other groups, suggest that permeability changes may result from changes in the cytoskeleton organization or from time-dependent local changes in the concentrations of ions in the vicinity of the channel, a phenomenon known as ion accumulation, that is at play in whole-cell experiments [156] and is further discussed below.

Li and collaborators have examined this carefully and have shown that P2X receptors readily activate and do not display a slow phase of activation under symmetrical ionic conditions [156]. These authors also show that changes in the equilibrium potential are observed with NMDG\(^+\) in the external solution during sustained activation and that this prolonged activation in bi-ionic solutions primes the depletion of internal Na\(^+\) and accumulation of NMDG\(^+\) in the cell. Li et al., also discuss that P2X receptors are exceptional in that they demonstrate significant permeability to large cations such as NMDG\(^+\) and that, even in the absence of pore dilation, would allow the conduction of relatively large molecules to enter or exit cells [156]. Finally, and most importantly, these authors point out that changes in ion concentration, produced by ion accumulation and revealed during experiments with P2X receptors, are an inherent problem in voltage-clamp recordings, where the flow of ionic currents across the membrane may alter the concentrations of intracellular ions [157,158]. In other words, when symmetrical ionic conditions are used, ion concentrations can be maintained by working under conditions in which current amplitudes are moderate and current measurements will always be equal to or greater than the flux of any individual ionic species. This is in contrast to experiments performed with asymmetric ionic conditions, which require a membrane conductance of an order of magnitude smaller than the access conductance to adequately control ion concentrations.

These results are prompting a reexamination of the concepts of pore dilation and dynamic ion selectivity for all ion channels [156], and highlight the importance of further studying the molecular mechanisms underlying different conformational changes induced by ligands in these proteins.

TRPV1 has been shown to contribute to several physiological and pathophysiological processes, importantly. Among these, the participation of TRPV1 in pain-related processes is an aspect that has attracted the attention of several research groups. Understanding cellular mechanisms leading to the generation of pain constitutes the basis for designing tools in order to attack them. Hence, research directed toward defining the molecular mechanisms that underlie the production of pain is of invaluable importance. This requires a great effort from many researchers and from several points of view, including that of understanding various particularities of the modulation of the effectors involved in pain processes.

As for changes in the conductance of TRPV1 due to the actions of agonists, it is interesting to note that, it has been recently reported that DkTx diminishes the unitary conductance by 32% of TRPV1, as compared to that observed with capsaicin [159]. DkTx is a molecule composed of two moieties joined by a short linker, which bind in the outer pore region of the channel so that each motif sits at a subunit interface [45]. Moreover, the effect was eliminated when the linker of DkTx was either absent or elongated and also when the pore turret of TRPV1 was removed. The lack of pore turret in the construct used for the cryo-EM of TRPV1 explains why the structure with DkTx/RTx is observed to be fully open. In fact, the unitary
conductance of the construct used to obtain the TRPV1 structure with bound DkTx was similar to that obtained with capsaicin [159].

With this, Geron et al. confirmed that DkTx and capsaicin elicit distinct gating mechanisms that are dependent upon the pore turret of TRPV1. They argued that the binding of DkTx directly restricts the movement of the pore helix and consequently the conductance.

In terms of the physiological relevance of LPA’s actions on the conductance of the TRPV1 channel, one can only hypothesize that in the presence of this agonist, as compared to capsaicin, depolarization of the neuron where it is expressed will occur more efficiently. Since the change in the conductance of the channel also occurs at subsaturating concentrations of LPA (i.e., 1 μM), this would mean that such an efficient depolarization could also occur even when lower levels of the phospholipid are present.

TRP channels share conserved sequences of amino acids in some regions of their structures. One such region is the TRP box (Figure 5), the region where the K710 residue that interacts with LPA lies. We have tested the effects of LPA on some TRP ion channels and have found that TRPV2, TRPV3, or TRPA1 channels are not activated by this phospholipid [41]. Although TRPV3 and TRPA1 have a lysine in this position, just as TRPV1, LPA is not capable of activating these channels. Thus, this phospholipid seems incapable of providing the energy necessary to drive the transition from the closed to the open state in this particular ion channel. This would further demonstrate the conformational versatility of this family of ion channels.

Several ion channels have been studied at great detail with respect to their responses to different agonists. In particular, the idea that LPA produces a conformational change that results in a different conductance of TRPV1 at the single-channel level compared to capsaicin challenges the notion in the field of study of ion channels in general.

Here we have discussed that LPA leads the TRPV1 ion channel to open with a single-channel conductance level that is higher than that attained with capsaicin. Both ligands promote nearly maximal open probabilities [117]; hence, this is most probably not a mechanism in which different subconductance levels can be attained in the presence of partial and full agonists.

Thus, it will be interesting to reflect upon and determine what mechanisms or structural rearrangements allow TRPV1, and maybe other TRP channels, to exhibit a higher conductance level in the presence of one agonist or another.

Acknowledgments

This work was supported by grants from Dirección General de Asuntos del Personal Académico (DGAPA)-Programa de Apoyo a Proyectos de Investigación e Innovación.
Disclosure statement
No potential conflict of interest was reported by the authors.

Funding
This work was supported by the Consejo Nacional de Ciencia y Tecnología [A1-S-8760]; Dirección General de Asuntos del Personal Académico, Universidad Nacional Autónoma de México [IN200717]; CONACYT A1-S-8760 to T.R.

References

[1] Hamill OP, Marty A, Neher E, et al. Improved patch-clamp techniques for high-resolution current recording from cells and cell-free membrane patches. Pflugers Arch. 1981;391(2):85–100.

[2] MacKinnon R. New insights into the structure and function of potassium channels. Curr Opin Neurobiol. 1991;1(1):14–19.

[3] Pore Loops: MR. An emerging theme in ion channel structure. Neuron. 1995;14(5):889–892.

[4] Potassium MR. Channels and the atomic basis of selective ion conduction (nobel Lecture). Angew Chem Int Ed Engl. 2004;43(33):4265–4277.

[5] Doyle DA, Morais Cabral J, Pfuetzner RA, et al. The structure of the potassium channel: molecular basis of K+ conduction and selectivity. Science. 1998;280(5360):69–77.

[6] Agre P. Aquaporin water channels (nobel lecture). Angew Chem Int Ed Engl. 2000;43(33):4278–4290.

[7] Yasui M, Kwon TH, Knepper MA, et al. Aquaporin-6: an intracellular vesicle water channel protein in renal epithelia. Proc Natl Acad Sci U S A. 1999;96(10):5808–5813.

[8] Cosens DJ, Manning A. Abnormal electroretinogram from a drosophila mutant. Nature. 1969;224(5216):285–287.

[9] Montell C, Rubin GM. Molecular characterization of the drosophila trp locus: A putative integral membrane protein required for phototransduction. Neuron. 1989;2(4):1313–1323.

[10] Hardie RC, Minke B. The Trp gene is essential for a light-activated Ca2+ channel in drosophila photoreceptors. Neuron. 1992;8(4):643–651.

[11] Caterina MJ, Schumacher MA, Tominaga M, et al. The cloned capsaicin receptor: A heat-activated ion channel in the pain pathway. Nature. 1997;389(6653):816–824.

[12] Tominaga M, Caterina MJ, Malmberg AB, et al. The cloned capsaicin receptor integrates multiple pain-producing stimuli. Neuron. 1998;21(3):531–543.

[13] Perozo E, Cortes DM, Cuello LG. Structural rearrangements underlying K+-channel activation gating. Science. 1999;285(5424):73–78.

[14] VanDongen AMJ. K channel gating by an affinity-switching selectivity filter. Proc Natl Acad Sci U S A. 2004;101(9):3248–3252.

[15] Schoppa NE, Sigworth FJ. Activation of shaker potassium channels ii. An activation gating model for wild-type and V2 mutant channels. J Gen Physiol. 1998;111(2):313–342.

[16] Zheng J, Sigworth FJ. Intermediate conductances during deactivation of heteromultimeric shaker potassium channels. J Gen Physiol. 1998;112(4):457–474.

[17] Bao H, Hakeem A, Henteleff M, et al. Voltage-insensitive gating after charge-neutralizing mutations in the S4 segment of shaker channels. J Gen Physiol. 1999;113(1):139–151.

[18] Yifrach O, MacKinnon R. Energetics of pore opening in a voltage-gated K(+) channel. Cell. 2002;111(2):231–239.

[19] Tranpani JG, Andalib P, Consiglio JF, et al. Control of single channel conductance in the outer vestibule of the Kv2.1 potassium channel. J Gen Physiol. 2006;128(2):231–246.

[20] Immke D, Korn SJ. Ion-ion interactions at the selectivity filter evidence from K(+)-dependent modulation of tetræethylammonium efficacy in Kv2.1 potassium channels. J Gen Physiol. 2000;115(4):509–518.

[21] Tang L, Gamal El-Din TM, Payandeh J, et al. Structural basis for Ca2+ Selectivity of a voltage-gated calcium channel. Nature. 2014;505(7481):56–61.

[22] Virginio C, MacKenzie A, Rassendren FA, et al. Pore dilation of neuronal P2X receptor channels. Nat Neurosci. 1999;2(4):315–321.

[23] Khakh BS, Lester HA. Dynamic selectivity filters in ion channels. Neuron. 1999;23(4):653–658.

[24] Linguaglia E, de Weille JR, Bassilana F, et al. A modulatory subunit of acid sensing ion channels in brain and dorsal root ganglion cells. J Biol Chem. 1997;272(47):29778–29783.

[25] de Weille JR, Bassilana F, Lazdunski M, et al. Identification, functional expression and chromosomal localisation of a sustained human proton-gated cation channel. FEBS Lett. 1998;433(3):257–260.

[26] Chung M-K, Güler AD, Caterina MJ. TRPV1 shows dynamic ionic selectivity during agonist stimulation. Nat Neurosci. 2008;11(5):555–564.
[27] Munns CH, Chung M-K, Sanchez YE, et al. Role of the outer pore domain in transient receptor potential vanilloid 1 (TRPV1) channel permeability to large cations. J Biol Chem. 2015;290(9):5707–5724.

[28] Brake AJ, Wagenbach MJ, Julius D. New structural motif for ligand-gated ion channels defined by an ionotropic ATP receptor. Nature. 1994;371(6497):519–523.

[29] Khakh BS, Egan TM. Contribution of transmembrane regions to ATP-gated P2X2 channel permeability dynamics. J Biol Chem. 2005;280(7):6118–6129.

[30] Di Virgilio F, Schmalzing G, Markwardt F. The elusive P2X7 macropore. Trends Cell Biol. 2018;28(5):392–404.

[31] Woo DH, Jung SJ, Zhu MH, et al. Direct activation of transient receptor potential vanilloid 1 (TRPV1) by diacylglycerol (DAG). Mol Pain. 2008;4:42.

[32] Wen H, Östman J, Bubb KJ, et al. 20-hydroxyeicosatetraenoic acid (20-HETE) is a novel activator of transient receptor potential vanilloid 1 (TRPV1) channel. J Biol Chem. 2012;287(17):13868–13876.

[33] Smart D, Gunthorpe MJ, Jerman JC, et al. The endogenous lipid anandamide is a full agonist at the human vanilloid receptor (HVR1). Br J Pharmacol. 2000;129(2):227–230.

[34] Moveahed P, Jönsson BAG, Birnir B, et al. Endogenous unsaturated C18 N-acylethanolamines are vanilloid receptor (TRPV1) agonists. J Biol Chem. 2005;280(46):38496–38504.

[35] Chu CJ, Huang SM, De Petrocellis L, et al. N-oleoyl-dopamine, a novel endogenous capsaicin-like lipid that produces hyperalgesia. J Biol Chem. 2003;278(16):13633–13639.

[36] Yoshida T, Inoue R, Morii T, et al. Nitric oxide activates TRP channels by cysteine s-nitrosylation. Nat Chem Biol. 2006;2(11):596–607.

[37] Yang F, Ma L, Cao X, et al. Divalent cations activate TRPV1 through promoting conformational change of the extracellular region. J Gen Physiol. 2014;143(1):91–103.

[38] Jara-Oseguera A, Bae C, Swartz KJ. An external sodium ion binding site controls allosteric gating in TRPV1 channels. eLife. 2016;5. DOI:10.7554/eLife.13356.

[39] Patacchini R, Santiocci P, Giuliani S, et al. Pharmacological investigation of hydrogen sulfide (H2S) contractile activity in rat detrusor muscle. Eur J Pharmacol. 2005;509(2–3):171–177.

[40] Bohlen CJ, Priet A, Zhou S, et al. A bivalent tarantula toxin activates the capsaicin receptor, TRPV1, by targeting the outer pore domain. Cell. 2010;141(5):834–845.

[41] Nieto-Posadas A, Picazo-Juárez G, Llorente I, et al. Lysophosphatidic acid directly activates TRPV1 through a C-terminal binding site. Nat Chem Biol. 2011;8(1):78–85.

[42] Moiseenkova-Bell VY, Stanciu LA, Serysheva II, et al. Structure of TRPV1 channel revealed by electron cryo-microscopy. Proc Natl Acad Sci U S A. 2008;105(21):7451–7455.

[43] Liao M, Cao E, Julius D, et al. Structure of the TRPV1 ion channel determined by electron cryo-microscopy. Nature. 2013;504(7478):107–112.

[44] Lishko PV, Procko E, Jin X, et al. The ankyrin repeats of TRPV1 bind multiple ligands and modulate channel sensitivity. Neuron. 2007;54(6):905–918.

[45] Cao E, Liao M, Cheng Y, et al. TRPV1 structures in distinct conformations reveal activation mechanisms. Nature. 2013;504(7478):113–118.

[46] Caterina MJ, Rosen TA, Tominaga M, et al. A capsaiacin-receptor homologue with a high threshold for noxious heat. Nature. 1999;398(6726):436–441.

[47] Shibasaki K. Physiological Significance of TRPV2 as a mechanosensor, thermosensor and lipid sensor. J Physiol Sci JPS. 2016;66(5):359–365.

[48] Zhang F, Hanson SM, Jara-Oseguera A, et al. Engineering vanilloid-sensitivity into the rat TRPV2 channel. eLife. 2016;5. DOI:10.7554/eLife.16409.

[49] Yang F, Vu S, Yarov-Yarovoy V, et al. Rational design and validation of a vanilloid-sensitive TRPV2 ion channel. Proc Natl Acad Sci U S A. 2016;113(26):E3657–3666.

[50] Zubičević L, Le S, Yang H, et al. Conformational Plasticity in the Selectivity Filter of the TRPV2 Ion Channel. Nat Struct Mol Biol. 2018;25(5):405–415.

[51] Zubičević L, Herzik MA, Chung BC, et al. Cryo-Electron Microscopy Structure of the TRPV2 Ion Channel. Nat Struct Mol Biol. 2016;23(2):180–186.

[52] Zhang F, Swartz KJ, Jara-Oseguera A. Conserved allosteric pathways for activation of TRPV3 revealed through engineering vanilloid-sensitivity. eLife. 2019;8. DOI:10.7554/eLife.42756.

[53] Zubičević L, Herzik MA, Wu M, et al. Conformational ensemble of the human TRPV3 ion channel. Nat Commun. 2018;9(1):4773.

[54] Singh AK, McGoldrick LL, Sobolevsky AI. Structure and gating mechanism of the transient receptor potential channel TRPV3. Nat Struct Mol Biol. 2018;25(9):805–813.

[55] Liedtke W, Choe Y, Marti-Renom MA, et al. Vanilloid receptor-related osmotically activated channel (VR-OAC), a candidate vertebrate osmoreceptor. Cell. 2000;103(3):525–535.

[56] Strotmann R, Harteneck C, Nunnenmacher K, et al. OTRPC4, a nonelective cation channel that confers sensitivity to extracellular osmolarity. Nat Cell Bio. 2000;2(10):695–702.

[57] Güler AD, Lee H, Iida T, et al. Heat-evoked activation of the ion channel, TRPV4. J Neurosci Off J Soc Neurosci. 2002;22(15):6408–6414. 20026679.
[58] Watanabe H, Davis JB, Smart D, et al. Activation of TRPV4 Channels (HVR-L2/MTRP12) by Phorbol Derivatives. J Biol Chem. 2002;277(16):13569–13577.

[59] Watanabe H, Vriens J, Prenen J, et al. Anandamide and arachidonic acid use epoxygenesatrienoic acids to activate TRPV4 channels. Nature. 2003;424 (6947):434–438.

[60] Vriens J, Watanabe H, Janssens A, et al. Cell swelling, heat, and chemical agonists use distinct pathways for the activation of the cation channel TRPV4. Proc Natl Acad Sci U S A. 2004;101(1):396–401.

[61] Deng Z, Paknejad N, Maksaev G, et al. Cryo-EM and X-ray structures of TRPV4 Reveal Insight into Ion Permeation and Gating Mechanisms. Nat Struct Mol Biol. 2018;25(3):252–260.

[62] Macpherson LJ, Geierstanger BH, Viswanath V, et al. The pungency of garlic: activation of TRPA1 and TRPV1 in response to allicin. Curr Biol CB. 2005;15(10):929–934.

[63] Bautista DM, Movahed P, Hinman A, et al. Pungent products from garlic activate the sensory ion channel TRPA1. Proc Natl Acad Sci U S A. 2005;102(34):12248–12252.

[64] Story GM, Peier AM, Reeve AJ, et al. ANKTM1, a TRP-like channel expressed in nociceptive neurons, is activated by cold temperatures. Cell. 2003;112(6):819–829.

[65] Viswanath V, Story GM, Peier AM, et al. Opposite thermosensor in fruitfly and mouse. Nature. 2003;423 (6942):822–823.

[66] Caspani O, Heppenstall PA. TRPA1 and cold transduction: an unresolved issue?. J Gen Physiol. 2009;133(3):245–249.

[67] Karashima Y, Talavera K, Everaerts W, et al. TRPA1 acts as a cold sensor in vitro and in vivo. Proc Natl Acad Sci U S A. 2009;106(4):1273–1278.

[68] Vandewauw I, De Clercq K, Mulier M, et al. A TRP channel trio mediates acute noxious heat sensing. Nature. 2018;555(7698):662–666.

[69] Bandell M, Story GM, Hwang SW, et al. Noxious cold ion channel TRPA1 is activated by pungent compounds and bradykinin. Neuron. 2004;41(6):849–857.

[70] Hinman A, Chuang -H-H, Bautista DM, et al. TRP channel activation by reversible covalent modification. Proc Natl Acad Sci U S A. 2006;103(51):19564–19568.

[71] Salazar H, Llorente I, Jara-Oseguera A, et al. A single n-terminal cysteine in trpv1 determines activation by pungent compounds from onion and garlic. Nat Neurosci. 2008;11(3):255–261.

[72] Salazar H, Jara-Oseguera A, Hernández-García E, et al. Structural determinants of gating in the TRPV1 channel. Nat Struct Mol Biol. 2009;16(7):704–710.

[73] Peyrot Des Gachons C, Uchida K, Bryant B, et al. Unusual pungency from extra-virgin olive oil is attributable to restricted spatial expression of the receptor of oleocanthal. J Neurosci Off J Soc Neurosci. 2011;31(3):999–1009.

[74] Laursen WJ, Bagriantsev SN, Gracheva EO. TRPA1 channels: chemical and temperature sensitivity. Curr Top Membr. 2014;74:89–112.

[75] Samanta A, Kiselas J, Pumroy RA, et al. Structural insights into the molecular mechanism of mouse TRPA1 Activation and inhibition. J Gen Physiol. 2018;150(5):751–762.

[76] Jordt S-E, Bautista DM, Chuang -H-H, et al. Mustard oils and cannabinoids excite sensory nerve fibres through the TRP channel ANKTM1. Nature. 2004;247(6971):260–265.

[77] Cavannaugh EJ, Simkin D, Kim D. Activation of transient receptor potential potential a1 channels by mustard oil, tetrahydrocannabinol and Ca2+ reveals different functional channel states. Neuroscience. 2008;154(4):1467–1476.

[78] Paulsen CE, Armache J-P, Gao Y, et al. Structure of the TRPA1 ion channel suggests regulatory mechanisms. Nature. 2015;525(7570):552.

[79] Martinez GQ, Gordon SE. Multimerization of homo sapiens TRPA1 ion channel cytoplasmic domains. PloS One. 2019;14(2):e0207835.

[80] Voets T, Janssens A, Prenen J, et al. Mg2+-dependent gating and strong inward rectification of the cation channel TRPV6. J Gen Physiol. 2003;121(3):245–260.

[81] Den Dekker E, Hoenderop JGJ, Nilius B, et al. The epithelial calcium channels, TRPV5 & TRPV6: from identification towards regulation. Cell Calcium. 2003;33(5–6):497–507.

[82] McGoldrick LL, Singh AK, Saotome K, et al. Opening of the human epithelial calcium channel TRPV6. Nature. 2018;553(7687):233–237.

[83] Kovalevskaya NV, Bokhovchuk FM, Vuister GW. The TRPV5/6 calcium channels contain multiple calmodulin binding sites with differential binding properties. J Struct Funct Genomics. 2012;13(2):91–100.

[84] Nilius B, Prenen J, Vennekens R, et al. Pharmacological modulation of monovalent cation currents through the epithelial Ca2+ channel ECaC1. Br J Pharmacol. 2001;134(3):453–462.

[85] van Goor MKC, Hoenderop JGJ, van der Wijst J. TRP channels in calcium homeostasis: from hormonal control to structure-function relationship of TRPV5 and TRPV6. Biochim Biophys Acta Mol Cell Res. 2017;1864(6):883–893.

[86] Hughes TET, Lodowski DT, Huynh KW, et al. Structural basis of TRPV5 channel inhibition by econazole revealed by Cryo-EM. Nat Struct Mol Biol. 2018;25(1):53–60.

[87] Hughes TET, Pumroy RA, Yazici AT, et al. Structural insights on TRPV5 gating by endogenous modulators. Nat Commun. 2018;9(1):4198.

[88] Hoenderop JGJ, Voets T, Hoefs S, et al. Homo- and heterotetrameric architecture of the epithelial Ca2+ channels TRPV5 and TRPV6. Embo J. 2003;22 (4):776–785.

[89] Ohsawa M, Miyabe Y, Katsu H, et al. Identification of the sensory nerve fiber responsible for lysophosphatidic acid-induced allodynia in mice. Neuroscience. 2013;247:65–74.
[90] Ueda H, Matsunaga H, Olaposi OI, et al. Chemical signature of neuropathic pain. Biochim Biophys Acta. 2013;1831(1):61–73.

[91] Inoue M, Rashid MH, Fujita R, et al. Initiation of neuropathic pain requires lysophosphatidic acid receptor signaling. Nat Med. 2004;10(7):712–718.

[92] Inoue M, Ma L, Aoki J, et al. Autotaxin, a synthetic enzyme of lysophosphatidic acid (LPA), mediates the induction of nerve-injured neuropathic pain. Mol Pain. 2008;4:6.

[93] Chemin J, Patel A, Duprat F, et al. Lysophosphatid Acid-Operated K+ Channels. J Biol Chem. 2005;280(6):4415–4421.

[94] Stirling L, Williams MR, Morrelli AD. Dual Roles for RHOA/RHOKinase in the regulated trafficking of a voltage-sensitive potassium channel. Mol Biol Cell. 2009;20(12):2991–3002.

[95] Kittaka H, Uchida K, Fukuta N, et al. Lysophosphatidic acid-induced itch is mediated by signalling of LPA5 receptor, phospholipase D and TRPA1/TRPV1. J Physiol. 2017;595(8):2681–2698.

[96] Hernández-Araiza I, Morales-Lázaro SL, Canul-Sánchez JA, et al. Role of lysophosphatidic acid in ion channel function and disease. J Neurophysiol. 2018;120(3):1198–1211.

[97] Czirják G, Tóth ZE, Enyedi P. The two-pore domain K+ channel, TRESK, is activated by the cytoplasmic calcium signal through calcineurin. J Biol Chem. 2004;279(18):18550–18558.

[98] Chuang HH, Prescott ED, Kong H, et al. Bradykinin potentiates bradykinin receptor activation and binding. J Biol Chem. 2004;279(18):18550–18558.

[99] Prescott ED, Julius D. A modular PIP2 binding site as a determinant of capsaicin receptor sensitivity. Neuron. 2010;68(4):597–607.

[100] Klein RM, Ufret-Vincenty CA, Hua L, et al. Determinants of molecular specificity in phosphoinositide regulation. Phosphatidylinositol (4,5)-Bisphosphate (PI(4,5)P2) is the endogenous lipid regulating TRPV1. J Biol Chem. 2008;283(38):26208–26216.

[101] Rosasco MG, Gordon SE, Bajjalieh SM. Characterization of the functional domains of a mammalian voltage-sensitive phosphatase. Biophys J. 2015;109(12):2480–2491.

[102] Senning EN, Collins MD, Stratievska A, et al. Regulation of TRPV1 ion channel by phosphoinositide (4,5)-bisphosphate: the role of membrane asymmetry. J Biol Chem. 2014;289(16):10999–11006.

[103] Stein AT, Ufret-Vincenty CA, Hua L, et al. Phosphoinositide 3-kinase binds to TRPV1 and mediates NGF-stimulated TRPV1 trafficking to the plasma membrane. J Gen Physiol. 2006;128(5):509–522.

[104] Stratievska A, Nelson S, Senning EN, et al. Reciprocal regulation among TRPV1 channels and phosphoinositide 3-kinase in response to nerve growth factor. eLife. 2018;7. DOI:10.7554/eLife.38869.

[105] Ufret-Vincenty CA, Klein RM, Collins MD, et al. Mechanism for phosphoinositide selectivity and activation of TRPV1 Ion Channels. J Gen Physiol. 2015;145(5):431–442.

[106] Ufret-Vincenty CA, Klein RM, Hua L, et al. Localization of the PIP2 Sensor of TRPV1 Ion Channels. J Biol Chem. 2011;286(11):9688–9698.

[107] Matta JA, Miyares RL, Ahern GP. TRPV1 Is a Novel Target for Omega-3 Polyunsaturated Fatty Acids. J Physiol. 2007;578(1Pt 2):397–411.

[108] Morales-Lázar SL, Llorente I, Sierra-Ramírez F, et al. Inhibition of TRPV1 channels by a naturally occurring omega-9 fatty acid reduces pain and itch. Nat Commun. 2016;7:13092.

[109] Picazo-Juárez G, Romero-Suárez S, Nieto-Posadas A, et al. Identification of a binding motif in the S5 helix that confers cholesterol sensitivity to the TRPV1 Ion Channel. J Biol Chem. 2011;286(24):24966–24976.

[110] Liu M, Huang W, Wu D, et al. TRPV1, but not P2X, requires cholesterol for its function and membrane expression in rat nociceptors. Eur J Neurosci. 2006;24(1):1–6.

[111] Szoke E, Börzsei R, Tóth DM, et al. Effect of lipid raft disruption on TRPV1 receptor activation of trigeminal sensory neurons and transfected cell line. Eur J Pharmacol. 2010;628(1–3):67–74.

[112] Telezhkin V, Reilly JM, Thomas AM, et al. Structural requirements of membrane phospholipids for M-Type potassium channel activation and binding. J Biol Chem. 2012;287(13):10001–10012.

[113] Jang Y, Lee MH, Lee J, et al. TRPM2 Mediates the Lysophosphatidic Acid-Induced Neurite Retraction in the Developing Brain. Pflugers Arch. 2014;466(10):1987–1998.

[114] Yung YC, Stoddard NC, Chun J, et al. Pathophysiology. J Lipid Res. 2014;55(7):1192–1214.

[115] Cao E, Cordero-Morales JF, Liu B, et al. TRPV1 channels are intrinsically heat sensitive and negatively regulated by phosphoinositide lipids. Neuron. 2013;77(4):667–679.

[116] Morales-Lázar SL, Serrano-Flores B, Llorente I, et al. Structural determinants of the transient receptor potential 1 (TRPV1) channel activation by phospholipid analogs. J Biol Chem. 2014;289(35):24079–24090.

[117] Canul-Sánchez JA, Hernández-Araiza I, Hernández-García E, et al. Different agonists induce distinct single-channel conductance states in TRPV1 channels. J Gen Physiol. 2018;150(12):1735–1746.

[118] Madej MG, Ziegler CM. Dawning of a new era in TRP channel structural biology by cryo-electron microscopy. Pflugers Arch. 2018;470(2):213–225.

[119] Duan J, Li J, Zeng B, et al. Structure of the mouse TRPC4 ion channel. Nat Commun. 2018;9(1):3102.

[120] Duan J, Li J, Chen G-L, et al. Cryo-EM structure of the receptor-activated TRPC5 ion channel at 2.9 angstrom
[121] Fan C, Choi W, Sun W, et al. Structure of the human lipid-gated cation channel TRPC3. eLife. 2018;7. DOI:10.7554/eLife.467969.

[122] Tang Q, Guo W, Zheng L, et al. Structure of the receptor-activated human TRPC6 and TRPC3 ion channels. Cell Res. 2018;28(7):746–755.

[123] Vinayagam D, Mager T, Apelbaum A, et al. Electron cryo-microscopy structure of the canonical TRPC4 ion channel. eLife. 2018;7. DOI:10.7554/eLife.36615.

[124] Li J, Zhang X, Song X, et al. The structure of TRPC ion channels. Cell Calcium. 2019;80:25–28.

[125] Thakur DP, Tian J, Jeon J, et al. Critical roles of Gi/o proteins and phospholipase C-Δ1 in the activation of receptor-operated TRPC4 channels. Proc Natl Acad Sci U S A. 2016;113(4):1092–1097.

[126] Kraft R, Harteneck C. The mammalian melastatin-related transient receptor potential cation channels: an overview. Pflugers Arch. 2005;451(1):204–211.

[127] Zhang Z, Tóth B, Szollosi A, et al. Structure of a TRPM2 channel in complex with Ca2+ explains unique gating regulation. eLife. 2018;7. DOI:10.7554/eLife.36409.

[128] Wang L, Fu T-M, Zhou Y, et al. Structures and gating mechanism of human TRPM2. Science. 2018;362:6421.

[129] Huang Y, Winkler PA, Sun W, et al. Architecture of the TRPM2 channel and its activation mechanism by ADP-ribose and calcium. Nature. 2018;562(7725):145–149.

[130] Duan J, Li Z, Li J, et al. Structure of full-length human TRPM4. Proc Natl Acad Sci U S A. 2018;115(10):2377–2382.

[131] Autzen HE, Myasnikov AG, Campbell MG, et al. Structure of the human TRPM4 ion channel in a lipid nanodisc. Science. 2018;359(6372):228–232.

[132] Winkler PA, Huang Y, Sun W, et al. Electron cryo-microscopy structure of a human TRPM4 channel. Nature. 2017;552(7684):200–204.

[133] Duan J, Li Z, Li J, et al. Structure of the mammalian TRPM7, a magnesium channel required during embryonic development. Proc Natl Acad Sci U S A. 2018;115(35):E8201–E8210.

[134] Yin Y, Le SC, Hsu AL, et al. Structural basis of cooling agent and lipid sensing by the cold-activated TRPM8 channel. Science. 2019;363(6430):eaav9334.

[135] Yin Y, Wu M, Zubcevic L, et al. Structure of the cold- and menthol-sensing ion channel TRPM8. Science. 2018;359(6372):237–241.

[136] Perraud AL, Fleig A, Dunn CA, et al. ADP-ribose gating of the calcium-permeable LTRPC2 channel revealed by nudix motif homology. Nature. 2001;411 (6837):595–599.

[137] Runnels LW, Yue L, Clapham DE. TRP-PLIK, a bifunctional protein with kinase and ion channel activities. Science. 2001;291(5506):1043–1047.

[138] Tóth B, Iordanov I, Csanády L. Putative channym activity of TRPM2 cation channel is unrelated to pore gating. Proc Natl Acad Sci U S A. 2014;111 (47):16949–16954.

[139] Iordanov I, Tóth B, Szollosi A, et al. Enzyme activity and selectivity filter stability of ancient TRPM2 channels were simultaneously lost in early vertebrates. eLife. 2019;8. DOI:10.7554/eLife.44556.

[140] Tan C-H, MaNaughton PA. The TRPM2 ion channel is required for sensitivity to warmth. Nature. 2016;536(7617):460–463.

[141] Csanády L, Töröcsik B. Four Ca2+ ions activate TRPM2 Channels by binding in deep crevices near the pore but intracellularly of the gate. J Gen Physiol. 2009;133(2):189–203.

[142] Launay P, Fleig A, Perraud AL, et al. TRPM4 is a Ca2+ -activated nonselective cation channel mediating cell membrane depolarization. Cell. 2002;109(3):397–407.

[143] Hofmann T, Chubanov V, Gudermann T, et al. TRPM5 is a voltage-modulated and Ga(2+)-activated monovalent selective cation channel. Curr Biol CB. 2003;13(13):1153–1158.

[144] Ulrich ND, Voets T, Prenen J, et al. Comparison of functional properties of the Ca2+-activated cation channels TRPM4 and TRPM5 from mice. Cell Calcium. 2005;37(3):267–278.

[145] Liu D, Liman ER. Intracellular Ca2+ and the phospholipid PIP2 regulate the taste transduction ion channel TRPM5. Proc Natl Acad Sci U S A. 2003;100 (25):15160–15165.

[146] Zhang Z, Okawa H, Wang Y, et al. Phosphatidylinositol 4,5-bisphosphate rescues TRPM4 channels from desensitization. J Biol Chem. 2005;280(47):39185–39192.

[147] Nilius B, Prenen J, Voets T, et al. Intracellular nucleotides and polyamines inhibit the Ca2+-activated cation channel TRPM4b. Pflugers Arch. 2004;448(1):70–75.

[148] Liman ER. The Ca2+-Activated TRP Channels: TRPM4 and TRPM5. In: Liedtke WB, Heller S, editors. TRP ion channel function in sensory transduction and cellular signaling cascades. Boca Raton (FL): CRC Press/Taylor & Francis; 2007:203–211.

[149] Guo J, She J, Zeng W, et al. Structures of the calcium-activated, non-selective cation channel TRPM4. Nature. 2017;552(7684):205–209.

[150] Krapivinsky G, Krapivinsky L, Manasian Y, et al. The TRPM7 channel is cleaved to release a cholinomodifying kinase. Cell. 2014;157(5):1061–1072.

[151] Zou Z-G, Rios FJ, Montezano AC, et al. TRPM7, Magnesium, and Signaling. Int J Mol Sci. 2019;20(8). DOI:10.3390/ijms20081877

[152] Numata T, Okada Y. Molecular determinants of sensitivity and conductivity of human TRPM7 to Mg2+ and Ca2+. Channels Austin Tex. 2008;2(4):283–286.
[153] Yudin Y, Rohacs T. Regulation of TRPM8 Channel Activity. Mol Cell Endocrinol. 2012;353(1–2):68–74.

[154] Rohács T, Lopes CMB, Michailidis I, et al. PI(4,5)P2 regulates the activation and desensitization of trpm8 channels through the TRP domain. Nat Neurosci. 2005;8(5):626–634.

[155] Bautista D, Julius D. Fire in the hole: pore dilation of the capsaicin receptor TRPV1. Nat Neurosci. 2008;11(5):528–529.

[156] Li M, Toombes GES, Silberberg SD, et al. Physical basis of apparent pore dilation of ATP-activated P2X receptor channels. Nat Neurosci. 2015;18(11):1577–1583.

[157] Mathias RT, Cohen IS, Oliva C. Limitations of the whole cell patch clamp technique in the control of intracellular concentrations. Biophys J. 1990;58(3):759–770.

[158] Frazier CJ, George EG, Jones SW. Apparent change in ion selectivity caused by changes in intracellular K(+) during whole-cell recording. Biophys J. 2000;78(4):1872–1880.

[159] Geron M, Kumar R, Zhou W, et al. TRPV1 pore turret dictates distinct DKTX and capsaicin gating. Proc Natl Acad Sci U S A. 2018;115(50):E11837–E11846.