Indole and other aromatic compounds activate the yeast TRPY1 channel

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Abstract The yeast TRPY1 (Yvc1p) channel is activated by membrane stretch to release vacuolar Ca\textsuperscript{2+} into the cytoplasm upon osmotic upshock. Exogenously added indole greatly enhances the upshock-induced Ca\textsuperscript{2+} release in vivo. Indole also reversibly activates the channels under patch clamp. A minimum of 10\textsuperscript{-6} M Ca\textsuperscript{2+} is needed for membrane stretch force to open TRPY1, but indole activation appears to be Ca\textsuperscript{2+} independent. A deletion of 30 residues at the predicted cytoplasmic domain, 570–600Δ, renders TRPY1 insensitive to stretch force up to 10\textsuperscript{-3} M Ca\textsuperscript{2+}. Nonetheless, indole readily activates this mutant channel. Several other aromatic compounds, e.g. the antimicrobial parabens, also activate TRPY1. These compounds likely alter the innate forces in the lipid bilayer received by the channel.

Keywords: TRP channel; Aromatic compound; Mechanosensitivity; Lipid bilayer; YVC1

1. Introduction

Aromatic compounds favor the two membrane–water interface regions of a lipid bilayer. For example, indole has this disposition, whether in its free form, or in the form of tryptophan as either a free amino acid [1], a peptide [2] or a protein [3]. Belts of aromatic residues [4], are found at the level of the interfaces [5] in many membrane proteins including ion channels KcsA [6], KirBac1.1 [7], etc.

TRPY1 (Yvc1p), in Saccharomyces cerevisiae, is a member of the “transient receptor potential” superfamily [8]. This channel corresponds to a 320-pS cation conductance in the yeast vacuolar membrane that can be activated by membrane stretch force under patch clamp [9]. It releases Ca\textsuperscript{2+} from the vacuole into the cytoplasm when live yeast cells are confronted with a sudden osmotic upshock [10]. TRPY1 activated by cytoplasmic Ca\textsuperscript{2+} provides a positive feedback in a Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release (CICR). We also isolated “gain-of-function” mutants that give larger Ca\textsuperscript{2+} signals at mild upshocks [11,9,12]. These mutant channels show kinetic defects, indicative of destabilization of various functional states (conformations) [11,12]. Seven of the ten gain-of-function mutations are in the predicted transmembrane domains of TRPY1. Surprisingly, they all involve aromatic amino-acid residues, leading to the conclusion that at least some of these aromatics are anchors for state stabilization [12]. In one case, the wild-type tyrosine was replaced with all 19 possible amino acids. Only replacements to the other two aromatic residues (trp and phe) retain the wild-type like channel kinetics [11]. We therefore decided to investigate whether exogenously added aromatic compounds can affect TRPY1 behavior.

2. Materials and methods

2.1. Yeast Strains and media
The wild-type parental strain, BY4742 (MAT\textalpha his3\Delta, leu2\Delta, lys2\Delta, ura3\Delta), and BY4742 yvc::kanmx4 (the yvc1Δ mutant) were used. Cells were cultured in eitherYPD (yeast extract peptone with glucose) for patch-clamp experiments cells or in CMD leu-minus (complete minimal medium with glucose, leucine dropout) for luminometry.

2.2. Luminometry
The transgenic-aequorin-based luminometric assay of Ca\textsuperscript{2+} release through TRPY1 was as described [11,12].

2.3. Patch clamp
Vacuolar preparation, patch clamp, and data acquisition and analyses were as described [8].

3. Results

3.1. Indole enhances upshock-induced Ca\textsuperscript{2+} release in vivo
Free Ca\textsuperscript{2+} in the cytoplasm of live yeast cells can be measured by the luminescence of transgenic aequorin [13]. A jump in the osmolarity of the BAPTA-containing medium causes an immediate rise of cytoplasmic Ca\textsuperscript{2+} in yeast cells, peaking in tens of second. This wave of Ca\textsuperscript{2+} is not observed in yvc1Δ, showing that it is the result of a release from the vacuole through the TRPY1 channel [10–12]. We found exogenously added indole to greatly enhance this response to osmotic upshock. This enhancement is robust and consistently observed (n > 60 experiments). Fig. 1, left, shows that 1 mM indole increases the peak response by more than 10 fold. There is no response to the upshock, with or without indole, when the YVC1 gene is deleted (Fig. 1, right).
3.2. Indole activates TRPY1 channel under patch clamp

Channel activation by indole was readily observed in either whole-vacuole mode (Fig. 2B) or in excised cytoplasmic-side-out mode of recording (Fig. 2C) ($n > 30$, activation observed in every preparation). Stretching the membrane with applied pressures up to lysis failed to activate the channel at or below $10^{-7}$ M Ca$^{2+}$ in repeated independent experiments ($n > 10$) (Fig. 2C, left two segments). Indole bypassed this requirement and readily activated channels at $10^{-5}$ M Ca$^{2+}$ (Fig. 2C, right-most segment) or even $10^{-6}$ M Ca$^{2+}$ (not shown). Once activated at $10^{-7}$ M Ca$^{2+}$ the channels were responsive to applied pressure. Submillimolar indole is rarely effective. Within the millimolar range, however, open probability rises steeply with concentration (Fig. 3A and B). Calculating directly from the concentration of indole in water and DMSO, the activation has a Hill coefficient of $\approx 6$, suggesting cooperativity. Unitary conductance when activated by indole is indistinguishable from that activated by Ca$^{2+}$ (Fig. 3C). Activated with Ca$^{2+}$ or stretch force, TRPY1 rectifies inwardly, shows a nonspecific, cation selectivity and $\approx 300$-pS unitary conductance when recorded in symmetric 150 mM K$^+$ [8,9]. All these properties are the same when channels are activated by indoles.

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Fig. 1. Indole potentiates up-shock-induced Ca$^{2+}$ release through TRPY1 (Yvc1p) in vivo. The luminescence of wild-type (left) or yvc1Δ (right) yeast cells expressing aequorin were examined. At the first arrow, nine volumes of either 0.5% DMSO (− indole curve) or 1 mM indole in 0.5% DMSO were added to the cells. Sorbitol was added at the second arrow at a final concentration of 1 M. Indole’s ability to potentiate the wild-type luminescent signal (in relative luminescence units, RLU) is evident (left). No response was seen in yvc1Δ cells (right) with the same indole treatment, indicating that the Ca$^{2+}$ release and the indole effect are entirely due to the TRPY1 channel.

Fig. 2. Indole activates the wild-type TRPY1. In whole-vacuole recording mode (A, B) and in excised cytoplasmic-side-out patch mode (C); currents flow into the cytoplasm are shown downward by convention. (A) Ca$^{2+}$ activation. Increasing [Ca$^{2+}$] from $10^{-7}$ M to $10^{-3}$ M Ca$^{2+}$ (arrow) activated an ensemble of the $\approx 300$-pS unitary conductances (in 150 mM KCl). (B) Indole activation. Addition of indole in 1% DMSO to the final concentration of 2 mM indole activated the ensemble current as well. C. Unitary conductances were activated in an excised patch with $10^{-5}$ M Ca$^{2+}$ (left segment). Activities subsided after perfusion of $10^{-7}$ M Ca$^{2+}$. At this [Ca$^{2+}$], the application of 120 mm Hg pressure failed to activate the channel (second segment). The same patch bathed with 1% DMSO control solution did not activate the channel (third segment), but the addition of 2 mM indole did (final segment).
3.3. Indole activates 570–600Δ

We have carried out extensive scanning and deletion analyses of the TPRY1 protein and the findings will be detailed elsewhere. Of interest here is a 30 amino acid deletion in the predicted cytoplasmic domain of TRPY1 (between position 570 and 600, diagramed at the top of Fig. 4). This produced a channel that can no longer be activated by high concentration of Ca²⁺ (10⁻³ M), membrane stretch force (120 mm Hg),

Fig. 4. 570–600Δ TRPY1 fails to respond to Ca²⁺ and applied stretch force but can be activated by indole. (Top trace) In a patch excised from a wild-type vacuole, two conducting units were inactive activated by 10⁻³ M Ca²⁺, and further activated by the exertion of 120 mm Hg pressure (left). Addition of 10⁻³ M Ca²⁺ activated the two units nearly completely, such that the additional effect of applied pressure was not evident (right). (Bottom trace) In a comparable experiment with a patch excised from a 570–600Δ vacuole, no channel activities could be elicited with 10⁻³ M Ca²⁺ and pressure (left), or even with 10⁻³ M Ca²⁺ and pressure (right). Addition of 2 mM indole clearly activated a unitary conductance in this patch.
or the combination of the two (Fig. 4, bottom trace, center segment). Nevertheless, indole readily activates the 570–600D channel (Fig. 4, last segment) in patches without BAPTA. Like the wild type, 570–600D channels are not active at 10^−7 M, but the addition of indole at this low [Ca^2+] readily activates them. Once activated these channels can be further activated by pressure (data not shown).

3.4. Several other aromatics also activate

To see whether the effects are unique to indole, we tested several other aromatic compounds. In repeated trials, activation was observed in both patch clamp and luminometry when indole was methylated in position 1, 2, 3, 5, or 7 of the indole ring (diagrammed in Fig. 5A). Quinoline, with two six-member rings, also activates. We also tested para-hydroxybenzoates, commonly called parabens. At millimolar concentrations we found that while para-hydroxybenzoic acid (paraben) did not activate, propylparaben activates TRPY1 in vivo and in vitro as effectively as indole. As shown in Fig. 5, 1 mM propylparaben enhances the upshock-induced release of Ca^{2+} into the cytoplasm of live yeast cells in a yvc1-dependent manner (Fig. 5B) similar to the effect of indole (Fig. 1). Under patch clamp, channel activation by propylparaben, even at 10^−7 M Ca^{2+}, is evident (Fig. 5C) (n = 4). Tryptamine, the amine equivalent of tryptophan, weakly activates. On the other hand, the comparable derivative of tyrosine, tyromine, activates very poorly; and that of phenylalanine, phenylamine, does not activate at all. In short, several of the very different aromatic compounds tested activate TRPY1, while others do not. This suggests that aromatic character maybe a key property but that there are additional contributing factors.

4. Discussion

How do these aromatic compounds activate TRPY1? One possibility is that the compound enters the interior of the channel protein to affect gating. The millimolar concentration requirement and steep dependency indicates that multiple low affinity binding sites are involved. Therefore, the mechanism is not likely to be as specific as those found with aromatic agonists and antagonists of dihydropyridine (DHP)-sensitive calcium channels [14]. Furthermore, the lack of geometrical specificity suggests that the site is not as restrictive as seen with indole inhibition of Shaker K^+ but not IRK currents [15]. In both these cases experimental evidence indicates that specific aromatic residues in the protein are important for the effect on channel activity. Our recent mutagenesis studies of TRPY1 show that more than one aromatic side chain plays a role in maintaining the closed and open form of the channel [11,12]. However, some aromatic compounds did not activate, implying that there are additional requirements that have yet to be clarified. One possible factor might be that the network of aromatic side chains in TRPY1 require compounds to penetrate to a certain depth into the interface in order to influence the side chains that play a role in maintaining the protein’s dynamic structure. We found that the compounds which activated tended to have greater calculated positive partitioning coefficients (log P or log D) into hydrophobic environments. Paraben has been shown to activate other channels including TRPA1 [16], a K^+ efflux in bacteria involving OmpF [17] and the mechanosensitive bacterial channels MscL and MscS [18]. In the latter case paraben was proposed to work from within the channel protein [18].
A second possibility is that the aromatic compounds act at the channel-lipid boundary. The high concentrations of aromatics needed to activate TRPY1 make this unlikely. L- and D-form of tarantula venom toxin GsMTx4 inhibits the stretch-activated channel (SACs) of vertebrate cells [19] by acting at this boundary. However, GsMX4 works at micromolar concentration to inhibit SACs [19], while aromatics work at millimolar to activate TRPY1 (present study) and TRPA1 [16]. If the exogenous aromatics compete with the channel protein's aromatic residue Y458 for lipid anchors, the un-anchored channel might be expected to show the “gain-of-function” phenotype, rapidly flickering between the closed and the open state as previously described [11,12]. However, as shown in Fig. 2B, “gain-of-function” flickers were not observed upon opening with aromatic compounds. Flickers were observed in prolonged incubations of some of patches, such that a boundary effect of indole cannot be completely ruled out at the later stage of activation.

It also seems likely that, at these concentrations, the compounds significantly affect the properties of the entire lipid bilayer, which then influences TRPY1 gating. Agents that alter bilayer elasticity are known to regulate the activities of gramicidin A and the voltage-dependent Na⁺ channel [20]. Calculations [21] and molecular dynamic simulations [22] show that there is a large tension in the bilayer at the two-water–membrane interfaces. Amphipathic compounds activate mechanosensitive channels such as MscS, MsCL [23], and TREK [24]. The general model is that entry of such amphipathic compounds induces curvature stress in the monolayers and differential insertion into the two monolayers causes bilayer bending, both perturbing the bilayer's internal forces to favor conformational changes of the embedded channels. We observed that certain amphipaths such as chlorpromazine can activate TRPY1 but others such as triitrophenol and lysophosphatidylincholine do not (unpublished results). The blockade of the natural activations by deletion 570–600 D-phosphotidylcholine do not (unpublished results). The blockage of the natural activations by deletion 570–600 D-phosphotidylcholine do not (unpublished results).

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