Identification of Inhibitory Ca\textsuperscript{2+} Binding Sites in the Upper Vestibule of the Yeast Vacuolar TRP Channel

Yeast vacuole cytosol TRPY1

TRPY1 current is inhibited by vacuolar Ca\textsuperscript{2+}

Vacular Ca\textsuperscript{2+}-mediated inhibition depends on D401 and D405

Ca\textsuperscript{2+} binds to D405 and D401 in the pore linker of TRPY1
Identification of Inhibitory Ca\(^{2+}\) Binding Sites in the Upper Vestibule of the Yeast Vacuolar TRP Channel

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SUMMARY

By vacuolar patch-clamp and Ca\(^{2+}\) imaging experiments, we show that the yeast vacuolar transient receptor potential (TRPY) channel 1 is activated by cytosolic Ca\(^{2+}\) and inhibited by Ca\(^{2+}\) from the vacuolar lumen. The channel is cooperatively affected by vacuolar Ca\(^{2+}\) (Hill coefficient, 1.5), suggesting that it may accommodate a Ca\(^{2+}\) receptor that can bind two calcium ions. Alanine scanning of six negatively charged amino acid residues in the transmembrane S5 and S6 linker, facing the vacuolar lumen, revealed that two aspartate residues, 401 and 405, are essential for current inhibition and direct binding of \(^{45}\text{Ca}^{2+}\). Expressed in HEK-293 cells, a significant fraction of TRPY1, present in the plasma membrane, retained its Ca\(^{2+}\) sensitivity. Based on these data and on homology with TRPV channels, we conclude that D401 and D405 are key residues within the vacuolar vestibule of the TRPY1 pore that decrease cation access or permeation after Ca\(^{2+}\) binding.

INTRODUCTION

The transient receptor potential (TRP) ion channel family is encoded by more than 100 genes (Venkatachalam and Montell, 2007). With the exception of the Ca\(^{2+}\)-selective TRPV5 and TRPV6 the TRP channels are non-selective cation channels sharing the membrane topology of six transmembrane helices (S1–S6) and N- and C-terminal domains residing within the cytosol. By mediating cation influx, TRP channels shape the membrane potential and increase the cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_{\text{cyt}}\)), translating environmental and endogenous stimuli into cellular signals. Most TRP channels fulfill their physiological function in the plasma membrane as cation influx channels, whereas some functional TRP channels are localized in the membrane of cytoplasmic organelles (Berbey et al., 2009; Chen et al., 2014; Dong et al., 2008; Lange et al., 2009; Oancea et al., 2009; Turner et al., 2003).

Calcium ions permeate TRP channels - with the exception of the Ca\(^{2+}\)-impermeable TRPM4 and TRPM5 - and also potentiate, activate, or inhibit currents most probably by interfering with TRP channel domains facing the cytosol (Blair et al., 2009; Bodding et al., 2003; Doerner et al., 2007; Du et al., 2009; Gross et al., 2009; Launay et al., 2002; McHugh et al., 2003; Olah et al., 2009; Prawitt et al., 2003; Starkus et al., 2007; Voets et al., 2002; Watanabe et al., 2003; Xiao et al., 2008; Zurborg et al., 2007). Likewise the TRP channel TRPY1, encoded by the vacuolar conductance 1 (YVC1) gene in Saccharomyces cerevisiae and localized in the vacuolar membrane, is activated by Ca\(^{2+}\) interfering with cytosolic channel domains (Bertl and Slayman, 1990; Palmer et al., 2001; Wada et al., 1987). TRPY1 mediates vacuolar Ca\(^{2+}\) release, and the amount of cytosolic [Ca\(^{2+}\)] required for TRPY1 activation can be substantially lowered in the presence of reducing agents such as dithiothreitol (DTT), glutathione, or \(\beta\)-mercaptoethanol (Bertl et al., 1992b; Bertl and Slayman, 1990). In addition to Ca\(^{2+}\), TRPY1 is also permeable to monovalent cations (permeability ratio P\(_{\text{K}^{+}}\)/P\(_{\text{K}^{+}}\) ~5; Bertl and Slayman, 1992), revealing a single-channel conductance of more than 300 pS in 180 mM KCl (Chang et al., 2010; Palmer et al., 2001; Wada et al., 1987; Zhou et al., 2003). TRPY1 is suggested to be involved in the response to hyperosmotic and oxidative stress as well as glucose-induced Ca\(^{2+}\) signaling (Bertl and Slayman, 1990; Bouillet et al., 2012; Denis and Cyert, 2002; Palmer et al., 2001). However, the exact function of the TRPY1-mediated Ca\(^{2+}\) release is still elusive.

Binding of Ca\(^{2+}\) by some mammalian TRP channels including the Ca\(^{2+}\)-activated TRPM4 has been described to be mediated by four coordinating residues within S2 and S3 close to the cytosolic S2-S3 linker (Autzen et al., 2018). These residues are not conserved in the TRPY1 sequence. All mammalian TRPs and the...
fly’s TRPL are binding Ca2+/calmodulin in vitro, and Bertl et al. (Bertl et al., 1992b, 1998a, 1998b) suggested that Ca2+/calmodulin contributes to TRPY1 activation in yeast, but a negative charge cluster, D573DDD576, within the TRPY1’s cytosolic C-terminus was shown to be crucial for the Ca2+-mediated activation of TRPY1 (Su et al., 2009).

In the present study, we characterized the dependence of TRPY1 current inhibition and activation on vacuolar and cytosolic [Ca2+] by patch-clamp recordings from yeast vacuoles and Ca2+ imaging in yeast. By alanine scanning and direct 45Ca2+ binding, we identified two aspartate residues within the S5-S6 linker facing the vacuolar lumen to be essential for Ca2+ binding and Ca2+-dependent inhibition of the TRPY1 current.

RESULTS

Cytosolic Ca2+ Activates the Vacuolar TRPY1 Channel

To get electrophysiological access to the yeast vacuole, the cell wall and membrane have to be removed (Figure 1A). At break-in and application of voltage ramps (150 to –150 mV), a small voltage-dependent outward rectifying current was recorded at membrane potentials above 80 mV (Figure 1B, bottom, blue current-voltage relation [IV]). Membrane potentials refer to the cytosolic side, and inward currents are defined as cation flow across the vacuolar membrane into the cytosol (Bertl et al., 1992a). Upon perfusion of the 150 mM KCl pipette solution, i.e., washout of vacuolar content, the outward current increases to maximum amplitude within 120 s (Figure 1B, red IV). Application of 1 mM cytosolic Ca2+ ([Ca2+]cyt) activates additional

Figure 1. Cytosolic Ca2+ Activates Cation Conductance in Yeast Vacuoles via TRPY1

(A) Procedure to release the yeast vacuole for patch-clamp experiments.

(B–E) Inward currents at –80 mV and outward currents at 80 mV, extracted from 200-ms ramps (0.5 Hz) spanning from 150 to –150 mV, Vh = 0 mV, plotted versus time (top), and corresponding current-voltage relations (IVs) at break-in (blue), after 120 s (red) and at maximum current (I_{max}) in 1 mM cytosolic Ca2+ (black; bottom) measured in isolated vacuoles from wild-type (WT; B, D, and E) and YVC1-deficient (YVC1 KO; C) yeast. The membrane potentials refer to the cytosolic side, i.e., inward currents represent movement of positive charges from the vacuole toward the cytosol (Bertl et al., 1992a). 1 mM Ca2+ was applied to the bath (representing the cytosol) as indicated by the bars. In (D) K+ and in (E) Cl– were substituted by TEA+ (tetraethylammonium) and gluconate in the patch pipette, respectively. Currents and IVs are shown as means ± SEM and just means, respectively, with the number of measured cells indicated in brackets.
inward and outward currents (Figure 1B, I_{max} in Ca), which are absent in vacuoles isolated from YVC1 knockout cells (YVC1 KO; Figure 1C). These currents are carried by cations, as their inward portion disappears in the absence of vacuolar (pipette) K+ substituted by TEA+ (tetraethylammonium) (Figure 1D). In contrast, the voltage-dependent outward rectifying current appearing after break-in is carried by Cl^-/CO_3^2-, as substitution of vacuolar Cl^-/CO_3^2- by gluconate significantly reduces its amplitude (Figure 1E, red IV). TRPY1 inward and outward cation currents were already detectable at [Ca^{2+}]_cyt of 10 µM (Figures 2A and 2B), and Ca^{2+} non-cooperatively (Hill coefficient < 1) increased currents in a concentration-dependent manner with apparent EC_{50} values of 498 µM for the inward current and 391 µM for the outward current (Figure 2B). 

Vacuolar Ca^{2+} Inhibits TRPY1 Activity

To characterize the dependence of TRPY1 currents on vacuolar [Ca^{2+}], we perfused the vacuole with 1 mM Ca^{2+} by the patch pipette. The currents activated by 1 mM [Ca^{2+}]_cyt were significantly reduced (Figures 2C and 2D). Figure 2C shows the vacuolar [Ca^{2+}]-dependent inhibition of TRPY1 currents activated by 1 mM [Ca^{2+}]_cyt with apparent IC_{50} values of 79 µM for the inward current and 109 µM for the outward current (Figure 2D). We consistently found Hill coefficients > 1 (1.4–1.5, Figure 2D) suggesting that the channel is inhibited by the cooperative binding of two molecules of Ca^{2+}. Inhibition of TRPY1 currents by vacuolar Ca^{2+} is still prominent at a pH value of 5.5 (Figure S1), considering the slightly acidic pH in the yeast vacuole (Preston et al., 1989).

Sr^{2+}, Ba^{2+}, and Mn^{2+} substitute for Ca^{2+} in TRPY1 activation (when present at the cytosolic site) and inhibition (when present at the vacuolar site) (Figure S2). In the absence of vacuolar Ca^{2+} ([Ca^{2+}]_vac), cytosolic concentrations of 1 mM divalent cations increased activation in the order Sr^{2+} (12%) < Mn^{2+} (15%) < Ba^{2+} (63%) with 100% current activation at 1 mM [Ca^{2+}]_cyt (Figures S2A and S2B). At 100% current activation (1 mM [Ca^{2+}]_cyt, and 0 [Ca^{2+}]_vac), vacuolar concentrations of 1 mM divalent cations inhibited in the order Mn^{2+} (33%) < Ba^{2+} (63%) < Sr^{2+} (68%) < Ca^{2+} (96%) (Figures S2C and S2D).
In HEK-293 cells TRPY1 retains its Ca2+-dependent properties

In yeast, TRPY1 was shown to be also activated by hyperosmotic shock, i.e., in the presence of high extra-cellular concentrations of sorbitol or NaCl (Zhou et al., 2003). Hyper- or hypoosmotic shock induces shrinkage or swelling of the vacuole, respectively, which easily disrupts the whole-vacuolar patch-clamp configuration. To prove whether hyperosmotic shock-induced TRPY1 activity is also inhibited by Ca2+, we expressed the YVC1 cDNA in HEK-293 cells. As shown by western blot the TRPY1 protein is present in transfected cells and a significant fraction is detectable in the plasma membrane by surface biotinylation (Figure 3A). In the plasma membrane the channel regions facing the vacuole in yeast are now facing the extracellular bath, whereas, like in yeast, the N-terminus, C-terminus, S2-S3, and S4-S5 linkers are facing the cytoplasm (Figure 3B). As shown in Figures 3C and 3D, TRPY1 currents are activated in the absence of extracellular Ca2+ by increasing the osmolarity in the bath by application of 0.5 and 0.75 M sorbitol, whereas no currents were detectable in non-transfected HEK-293 cells (Figure 3E). Applying 1 mM Ca2+ to the bath significantly reduced the TRPY1 current induced by 0.75 M sorbitol (Figure 3F, red trace) indicating that TRPY1 integrates activating and inhibitory stimuli also in HEK-293 cells.

Figure 3. Ca2+-Dependent Regulation of TRPY1 Currents in Transfected HEK-293 Cells

(A) Western blot of protein lysates from YVC1 cDNA-transfected (HEK TRPY1) and non-transfected (HEK) HEK-293 cells. Biotinylation of surface proteins shows that a considerable fraction of TRPY1 present in transfected HEK-293 cells (input) resides in the plasma membrane (biotinylated). The filter was stripped and incubated in the presence of an antibody directed against the intracellular protein calnexin as control. (B) Orientation of TRPY1 (red) in the plasma membrane with N- and C-termini residing within the cytosol. (C–K) Inward and outward currents at −80 and 80 mV, extracted from 400-ms ramps (0.5 Hz) spanning from −100 to 100 mV, Vh = 0 mV, plotted versus time (C, E–G, and I–K) and corresponding IVs (D and H) of maximum currents (Imax) from (C) or at 240 s from (G), activated by 0.5 or 0.75 M sorbitol (hyperosmotic shock; C–F) or indicated [Ca2+]cyt (int Ca; buffered with BAPTA; G–K) at 0 or 1 mM external (ext) Ca2+ in HEK TRPY1 cells (HEK TRPY1; C, D, F–H, J, and K) or non-transfected HEK-293 cells (HEK; E and I). The bars indicate application of 0.5 or 0.75 M sorbitol in C, E, and F, or 0 and 1 mM Ca2+-containing bath solution in K and J, respectively. The black traces in C and F are the same. Currents and IVs are normalized to the cell size (pA/pF) and are shown as means ± SEM (C, E–G, and I–K) and means (D and H) with the number of measured cells in brackets.

In HEK-293 Cells TRPY1 Retains Its Ca2+-Dependent Properties

In yeast, TRPY1 was shown to be also activated by hyperosmotic shock, i.e., in the presence of high extracellular concentrations of sorbitol or NaCl (Zhou et al., 2003). Hyper- or hypoosmotic shock induces shrinkage or swelling of the vacuole, respectively, which easily disrupts the whole-vacuolar patch-clamp configuration. To prove whether hyperosmotic shock-induced TRPY1 activity is also inhibited by Ca2+, we expressed the YVC1 cDNA in HEK-293 cells. As shown by western blot the TRPY1 protein is present in transfected cells and a significant fraction is detectable in the plasma membrane by surface biotinylation (Figure 3A). In the plasma membrane the channel regions facing the vacuole in yeast are now facing the extracellular bath, whereas, like in yeast, the N-terminus, C-terminus, S2-S3, and S4-S5 linkers are facing the cytoplasm (Figure 3B). As shown in Figures 3C and 3D, TRPY1 currents are activated in the absence of extracellular Ca2+ by increasing the osmolarity in the bath by application of 0.5 and 0.75 M sorbitol, whereas no currents were detectable in non-transfected HEK-293 cells (Figure 3E). Applying 1 mM Ca2+ to the bath significantly reduced the TRPY1 current induced by 0.75 M sorbitol (Figure 3F, red trace) indicating that TRPY1 integrates activating and inhibitory stimuli also in HEK-293 cells.
In the absence of extracellular Ca²⁺, TRPY1 inward and outward currents in HEK-293 cells are activated by [Ca²⁺]ₒᵣᵣ (at 3 and 10 μM; Figures 3G and 3H), confirming the assumed orientation of TRPY1 in the plasma membrane. Non-transfected HEK-293 cells do not reveal any significant currents under these conditions (Figure 3i). TRPY1 currents activated by 10 μM [Ca²⁺]ₒᵣᵣ in the absence of extracellular Ca²⁺ are readily inhibited when 1 mM Ca²⁺ was added (Figure 3j). Figure 3k shows that in the presence of 1 mM external Ca²⁺, almost no TRPY1 current appeared at [Ca²⁺]ₒᵣᵣ of 10 μM, whereas after removal of external Ca²⁺ a significant current developed (Figure 3k).

Aspartate Residues 401 and 405 Mediate the Ca²⁺-Dependent Inhibition of TRPY1

The mechanism of the inhibition of TRPY1 currents by vacuolar Ca²⁺ in yeast or extracellular Ca²⁺ in HEK-293 cells was not known. We noticed six negatively charged acidic residues in the S5–S6 linker contributing to the TRPY1 pore, D398, D401, D405, D425, E428, and E429 (Figure 4a), which we replaced by alanine residues. The cDNAs of all mutants and of wild-type TRPY1 (plasmids see Table 1) were expressed in YVC1 KO yeast (Chang et al., 2010). Wild-type and mutant proteins are detectable in western blot (Figure 4b). Only the transformation of yeast with plasmid pRS316-TRPY1D405A did not yield any colonies. Figures 4d–4h show currents of wild-type TRPY1 (Figure 4d) and of the TRPY1 mutants, expressed in YVC1 KO yeast (Figures 4e–4h), activated by 1 mM [Ca²⁺]ₒᵣᵣ in the absence (Ca₀) or presence of 1 mM vacuolar Ca²⁺. As shown for vacuoles of wild-type yeast cells (Figure 2) 1 mM [Ca²⁺]ₒᵣᵣ activates TRPY1 in the absence (Figure 4d, black trace), but not in the presence of vacuolar Ca²⁺ (Figure 4d, red trace), whereas TRPY1D398A failed to yield currents even in the absence of vacuolar Ca²⁺ (Figure 4e). The Ca²⁺-mediated outward current detectable in the absence of vacuolar Ca²⁺ most probably reflects changes of the vacuolar chloride conductance (see Figure 1). Expression of TRPY1D401A yielded a small constitutive current (Figure 3a, green trace), which could be significantly enhanced by 1 mM [Ca²⁺]ₒᵣᵣ, leading to current amplitudes more than five times higher than mediated by wild-type TRPY1 (Figure 4f, black trace), but currents were no longer inhibited by vacuolar Ca²⁺ (Figure 4f, red trace). Beside smaller current amplitudes, TRPY1D425A and TRPY1E428A,E429A revealed the same Ca²⁺ dependence as TRPY1 wild-type (Figures 4g and 4h). Ca²⁺ imaging experiments obtained after expressing the cytosolic luminescent Ca²⁺ reporter ae–quorin in intact yeast confirmed the patch-clamp data (Figure 4i). TRPY1D401A induced the highest cytosolic Ca²⁺ increase upon hyperosmotic shock, whereas TRPY1D398A did not respond at all. Current-voltage relationships and statistics of whole-vacuole current amplitudes and yeast cytosolic Ca²⁺ signals are summarized in Figure S3. The reversal potential of vacuolar currents did not vary between TRPY1 mutants and wild-type, arguing against changes of ion permeability.

Wild-type and mutant YVC1-transformed YVC1 KO yeast showed different amplitudes of vacuolar currents and cytosolic Ca²⁺ signals: D401A revealed larger currents and higher [Ca²⁺]ₒᵣᵣ than wild-type TRPY1, whereas for TRPY1D425A and TRPY1E428A,E429A, currents and changes of [Ca²⁺]ₒᵣᵣ were smaller (Figures 4d–4i). Western blots revealed similar amounts of TRPY1 protein in yeast transformed with wild-type, D398A, D401A, and D425A YVC1, relative to the endogenous SRP1, used as a control (Figure 4c). The higher amount of TRPY1E428A,E429A protein (Figure 4c) not mirrored by larger currents (Figure 4h) and higher [Ca²⁺]ₒᵣᵣ (Figure 4i) might indicate that only part of the protein detectable in western blot is targeted to the vacuolar membrane.

Transformation of yeast with pRS316-YVC1D405A did not yield any colonies. To yield lower expression levels to prevent delirious effects of overexpression, we cloned the YVC1 gene (wild-type, YVC1D401A, and YVC1D405A) in single-copy pRS316 CEN plasmids without its own promoter. Under these conditions, cytosolic Ca²⁺-induced vacuolar currents were recorded from YVC1 wild-type and YVC1D401A- and YVC1D405A-transformed YVC1 KO yeast (Figures S4a–S4c). In contrast to wild-type TRPY1 (Figure S4a), the currents induced by cytosolic Ca²⁺ remained in the presence of 1 mM vacuolar Ca²⁺ in TRPY1D401A (Figure S4b; note the spontaneous current after break-in) and TRPY1D405A (Figure S4c).

Beside a smaller current amplitude of TRPY1D405A at 40 mV in the absence of vacuolar 1 mM Ca²⁺ (see Figure S4e), neither the mutations D401A and D405A nor the presence of vacuolar 1 mM Ca²⁺ significantly affected single-channel current amplitudes of TRPY1 analyzed at −40 and 40 mV (Figures 4j and S4e, respectively; example traces are shown in Figure S4d). Single-channel current amplitudes of TRPY1D405A were analyzed from yeast cells transfected with single-copy pRS316 CEN plasmids without its own promoter (see above). The unchanged single-channel current amplitude (Figures 4j and S4e) but significantly

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See also Figures S3 and S4.

indicated. One-way analysis of variance (ANOVA) revealed no differences.

including TRPY1D405A were detectable (Figure 5A), and we recorded hyperosmotic shock (Figures 5B–5G) transformation (x/y; (I) indicated in brackets. Single-channel current amplitudes in (J) are shown as means

Figure 4. Vacuolar TRPY1 Currents in YVC1 KO Yeast Expressing Wild-Type or Mutant YVC1 cDNAs

(A) Predicted transmembrane topology of the yeast vacuolar TRPY1 (675 amino acid residues) with negative charge cluster at the cytosolic C-terminus assumed to be responsible for TRPY1 activation (blue) (Su et al., 2009) and negatively charged residues within the SS-S6 linker facing the vacuolar lumen (red). The TRPY1G402S mutant (green) is constitutively active (Zhou et al., 2007), and the TRPY1C624S mutant (black) can no longer be activated by 2-mercaptoethanol (Hamamoto et al., 2018).

(B) Western blot of protein lysates from non-transformed yeast (YVC1 KO) or YVC1 KO yeast transformed with wild-type (WT) and mutant YVC1 cDNAs (as indicated). Filter was stripped and incubated with antibodies for aequorin (middle) and SRP1 (serine-rich protein; bottom) as controls.

(C) Antibody stain intensities of TRPY1 proteins in (B) were quantified relative to the antibody stain of the endogenous serine-rich protein 1 (SRP1) and normalized to the WT TRPY1/SRP1 ratio (summary of five independent western blots). One-way analysis of variance (ANOVA): not significant (ns), *** p<0.001 compared to WT.

(D–H) Whole-vacuole currents at –80 and 80 mV extracted from 200-ms ramps (0.5 Hz) spanning from 150 to –150 mV, Vh = 0 mV, plotted versus time, activated by cytosolic Ca2+ (1 mM) at 0 (black) or 1 mM (red) vacuolar (vac, patch pipette) Ca2+ in YVC1 KO cells expressing WT (D) or mutant (E–H) YVC1 cDNAs. (I) Changes of cytosolic Ca2+ challenged by 1.5 M NaCl (hyperosmotic shock) monitored in transformed YVC1 KO yeast cells (as in D–H) as relative luminescence units (RLU). Bars indicate application of 1 mM Ca2+ (D–H) or 1.5 M NaCl (I).

(J) Single-channel current amplitudes of TRPY1WT, TRPYD401A, and TRPYD425A in the absence and presence of vacuolar 1 mM Ca2+*, analyzed at –40 mV from current traces of voltage ramps and voltage steps in whole vacuoles and excised (outside-out) vacuolar patches. Single channels of TRPY1D405A were analyzed from yeast cells transfected with single-copy pRS316 CEN plasmids without its own promoter (see Figure S4C). Currents are normalized to the size of the vacuole (pA/pF) and shown as means ± SEM with number of measured vacuoles (D–H) or the number of experiments (x) from y independent measurements (x/y; (I) indicated in brackets. Single-channel current amplitudes in (J) are shown as means ± SEM with number of independent experiments indicated. One-way analysis of variance (ANOVA) revealed no differences. See also Figures S3 and S4.

reduced whole-vacuolar current amplitude (Figure 4D) in wild-type TRPY1 suggests a prominent decrease of the open probability in the presence of vacuolar 1 mM Ca2+.

After expression of wild-type and mutant YVC1 cDNAs (plasmids see Table 1) in HEK-293 cells, all proteins including TRPY1D405A were detectable (Figure 5A), and we recorded hyperosmotic shock (Figures 5B–5G) as well as cytosolic Ca2+*-induced currents (Figures 5H–5J). Current-voltage relationships and statistics of
whole-cell current amplitudes are summarized in Figure S5. Compared with wild-type TRPY1 (Figure 5B) hyperosmotic shock did not initiate TRPY1D398A currents (Figure 5C), but the TRPY1D425A (Figure 5F) and TRPY1E428A,E429A mutants (Figure 5G) behaved like wild-type TRPY1, very similar as observed in the yeast vacuole (Figure 4). Like in yeast vacuoles TRPY1D401A yielded spontaneous currents (Figures 5D, 5E, and S5A, green trace), and, compared with wild-type TRPY1 (Figures 5B and 5H), extracellular Ca2+ (1 and 2 mM, respectively) did not inhibit hyperosmotic shock and cytosolic Ca2+-induced activity of the mutant channels TRPY1D401A and TRPY1D405A (Figures 5D, 5I, 5E, and 5J). The latter mutant induced even larger current amplitudes upon hyperosmotic shock in the presence of extracellular Ca2+. Current-voltage relationships and statistics of current amplitudes are summarized in Figure S5.

To prove whether residues D401 and D405 are required for Ca2+ binding 25-amino-acid (25-mer) peptides derived from wild-type (amino acid residues F390 to L414) and D401A and D405A mutant SS-S6 linkers of TRPY1 were synthesized and spotted onto hardened cellulose membranes and incubated with 45Ca2+.
(Figure 6). The wild-type peptide significantly binds $^{45}$Ca$^{2+}$, whereas the TRPY1-D401A and TRPY1-D405A peptides showed less and the double mutant (TRPY1-D401A,D405A) no detectable $^{45}$Ca$^{2+}$ binding. As control, we spotted a 25-mer peptide representing the first EF hand (EF1) of mammalian calmodulin (CaM; Figure 6). The $^{45}$Ca$^{2+}$ binding by EF1 was completely abolished when its glutamate residues were replaced by alanine residues (Figure 6). The amounts of $^{45}$Ca$^{2+}$ bound by EF1 and TRPY1-D401-D405 are very similar, as estimated by the signals’ intensities. Considering equal amounts of spotted 25-mer peptides (about 16 nmoles per spot), additionally estimated by UV absorption at 312 nm (Figure 6), close apparent binding affinities of the peptides for Ca$^{2+}$ might be assumed. At physiological salt concentration, the non-cooperative Ca$^{2+}$ binding affinity of the isolated CaM EF1 was estimated to be about 125 $\text{mM}$ (Ye et al., 2005), which is very close to the value of 79–109 $\text{mM}$ measured here for vacuolar Ca$^{2+}$-dependent inhibition of TRPY1 (Figure 2D).

**DISCUSSION**

We performed whole-vacuole patch-clamp recordings and Ca$^{2+}$ imaging experiments in yeast to study the Ca$^{2+}$ dependence of TRPY1 using as controls a YVC1-deficient yeast strain (Chang et al., 2010) and HEK-293 cells heterologously expressing the YVC1 cDNA as functional TRPY1 channels in the plasma membrane. We show that inhibition of vacuolar TRPY1 is cooperatively mediated by the vacuolar [Ca$^{2+}$] (IC$\text{_{50}}$ 79–109 $\mu$M), that it requires two aspartate residues (D401 and D405) in the S5-S6 linker facing the vacuolar lumen, and that these aspartate residues are directly involved in Ca$^{2+}$ binding. The yeast vacuolar TRP cation channel TRPY1 may be assumed as a progenitor of TRP channels present in mammals, flies, and worms and underlies cation movements across the yeast vacuolar membrane. Additional TRP-related proteins in *Saccharomyces cerevisiae* like FLC2, encoded by the YAL053W gene, seem to be involved in endoplasmic reticulum Ca$^{2+}$ release (Rigamonti et al., 2015). TRPY1 is activated by osmotic stress (Batiza et al., 1996; Denis and Cyert, 2002; Zhou et al., 2003), reducing agents (Bertl and Slayman, 1990, Wada et al., 1987), aromatic compounds like indole (John Haynes et al., 2008), and cytosolic Ca$^{2+}$ (Bertl and Slayman, 1990, Wada et al., 1987). In the presence of DTT the EC$\text{_{50}}$ values of TRPY1 activation by cytosolic Ca$^{2+}$ are 498 $\mu$M (inward current) and 391 $\mu$M (outward current) with Hill coefficients < 1. Reducing agents such as DTT or β-mercaptoethanol lower the [Ca$^{2+}$] required for channel activation (Bertl and Slayman, 1990, 1992), and the cysteine residue 624 within TRPY1’s C terminus facing the cytosol (Figure 4A) has been suggested to be involved in this process (Hamamoto et al., 2018). Reducing agents may mimic the redox environment in the yeast cytosol (Carpaneto et al., 1999; Lopez-Mirabal and Winther, 2008; Palmer et al., 2001), which might contribute to TRPY1 activation under certain conditions. Vacular TRPY1 currents are already activated at [Ca$^{2+}$] of 10 $\mu$M of cytosolic Ca$^{2+}$ (Figure 2 and Bertl et al., 1992b; Bertl and Slayman, 1990, 1992), but the yeast [Ca$^{2+}$] has been estimated to be in the range of 260 nM (Halachmi and Eilam, 1989). Probably, hyperosmotic shock represents the initial physiological stimulus in intact yeast, which increases cytosolic Ca$^{2+}$, which in turn facilitates TRPY1 activity. Palmer et al. (Palmer et al., 2001) suggested that the cytosolic activation of TRPY1 is Ca$^{2+}$ specific, but we also detected activation by the alkaline earths Ba$^{2+}$ > Sr$^{2+}$ and by the transition metal ion Mn$^{2+}$ (Figure S2).
High $[\text{Ca}^{2+}]$ within the vacuolar lumen presumably inhibits TRPY1 currents (Zhou et al., 2003), and 1 mM Ca$^{2+}$ added to the bath eliminates TRPY1 single-channel currents in inside-out excised vacuolar patch-clamp recordings (Hamamoto et al., 2018). The present study shows that vacuolar Ca$^{2+}$, as well as Sr$^{2+}$ and Ba$^{2+}$, strongly inhibit TRPY1 activity and that the amino acid residues D401 and D405 located in the vacuolar loop between S5 and S6 (Figure 4A) are crucial for Ca$^{2+}$-dependent inhibition and direct Ca$^{2+}$ binding. Mutation of the nearby G402 (Figure 4A) renders TRPY1 constitutively active (Zhou et al., 2007), indicating the significance of this area for proper gating. Hamamoto et al. described that luminal Zn$^{2+}$ increased TRPY1 currents in vacuolar excised patches (Hamamoto et al., 2018). Zn$^{2+}$ might be attracted to a single negatively charged residue and thereby prevent efficient coordination of Ca$^{2+}$. In mouse TRPV3 and TRPV4, corresponding residues D641 and D682, respectively, were shown to be key residues for extracellular Ca$^{2+}$-mediated channel inhibition (Voets et al., 2002; Watanabe et al., 2003; Xiao et al., 2008), and human TRPV1 D646 was suggested to be a high-affinity binding site for cations (Garcia-Martinez et al., 2000). However, D646 and D682 are part of the selectivity filter in TRPV1 and TRPV4, respectively (Deng et al., 2018; Liao et al., 2013), and, by sequence comparison, rather align to TRPY1D425 (Figure S6), whereas TRPY1D401 and TRPY1D405 are located in the vacuolar vestibule of TRPY1’s ion-conducting pore. The residues E600, D601, E610, and E648 are involved in pH-dependent modulation of human TRPV1 (Jordt et al., 2000), with TRPY1D401 aligning to TRPY1D425 (Figure S6). Although nothing is known about external Ca$^{2+}$-mediated inhibition of TRPV1 and TRPV6, their S5-S6 linkers appear closer related to TRPY1 than TRPV3, TRPV4, TRPV2, and TRPA1 by sequence alignment (Figure S6). TRPV6 residues D517 and E518, located in the upper vestibule and aligning with TRPY1D401, are shown to bind and guide Ca$^{2+}$ into the channel pore of TRPV6 (Singh et al., 2017).

The structure of TRPY1 is not yet available. Therefore, we aligned the structure of TRPY1 according to the published structure of Xenopus tropicalis TRPV4 (Deng et al., 2018), which is inhibited by Ca$^{2+}$ from the extracellular side (see above; Figures S6B and S6C). Our data show that TRPY1 D401 and D405 are decisive for the Ca$^{2+}$-dependent inhibition of the channel. Both residues, D401 and D405, also highly conserved in TRPY homologs of other fungi (Ihara et al., 2014), are part of the outer pore domain of the channel and orienated to the lumen of the vacuole and are directly involved in Ca$^{2+}$ binding. The alignment suggests that binding of Ca$^{2+}$ to residues D401 and D405 might result in a reorientation of

### Table 1. TRPY1 Plasmids Generated in the Laboratories of the Authors and Used in This Study

| Plasmid                          | TRPY1 Construct                  |
|---------------------------------|----------------------------------|
| Backbone: Yeast single-copy centromeric pRS316 vector |                        |
| pGS2062                         | TRPY1 wild-type                  |
| pSB276                          | TRPY1D398A                       |
| pSB277                          | TRPY1D401A                       |
| pSB278                          | TRPY1D405A                       |
| pSB279                          | TRPY1D425A                       |
| pSB280                          | TRPY1E428A,E429A                 |
| Backbone: Bicistronic eukaryotic expression (HEK-293 cells) |                        |
| pCAGGS-YVC1-IRES-GFP            | TRPY1 wild-type                  |
| pCAGGS-YVC1D398A-IRES-GFP       | TRPY1D398A                       |
| pCAGGS-YVC1D401A-IRES-GFP       | TRPY1D401A                       |
| pCAGGS-YVC1D405A-IRES-GFP       | TRPY1D405A                       |
| pCAGGS-YVC1D425A-IRES-GFP       | TRPY1D425A                       |
| pCAGGS-YVC1E428A,E429A-IRES-GFP | TRPY1E428A,E429A                 |
residues I455 and Y458, possibly involved in forming the lower gate of TRPY1, and thereby narrowing or occluding the lower gate (Figure S6B). According to the model, replacing D401 and D405 by alanine residues disables Ca2+ binding and thus prevents such structural changes in the presence of vacuolar Ca2+ (Figure S6C). As single-channel current amplitudes of TRPY1 are not affected (Figure 4J), we assume that Ca2+ binding to D401 and D405 affects the open probability by rearranging the main S6 segment, stabilizing a closed inactivated conformation of the channel by narrowing and occluding the lower gate.

The physiological pH value in yeast vacuoles is slightly acidic (Pearce et al., 1999; Preston et al., 1989). Vacuolar pH 6.0 does not alter the mechanosensitivity of TRPY1 (Zhou et al., 2003), whereas the current activated by cytosolic Ca2+ was reduced at a vacuolar pH 5.5, but the vacuolar Ca2+-dependent inhibition remained (Figure S1). The mammalian TRP channels TRPML1, 2, and 3 are located in endolysosomes, and thus exert their functions also in a cytoplasmic organelle environment as TRPY1. TRPML channels are involved in the release of Ca2+ and Fe2+ into the cytosol (Cheng et al., 2010), and TRPML1 has been shown to be regulated by lysosomal Ca2+ and pH (Cantiello et al., 2005; Li et al., 2017). Like for wild-type TRPY1 a significant fraction of the constitutively active TRPML1 mutant, TRPML1V432P, can be expressed in the plasma membrane of HEK-293 cells. Extracellular Ca2+ inhibits TRPML1V432P channel activity with an apparent IC50 of 270 μM (Li et al., 2017), which is in the range of the vacuolar [Ca2+], which inhibits TRPY1 (Figure 2). External (lysosomal) acidification reduced the Ca2+-dependent inhibition of TRPML1V432P, and an electronegative luminal pore domain was indicated as a unique hallmark of TRPML1’s interaction site for lysosomal Ca2+ and H+ (Li et al., 2017).

The yeast vacuolar Ca2+ concentration was estimated to be 1.3 mM (Halachmi and Eilam, 1989; Iida et al., 1990) and the vacuole was assumed to function as a Ca2+ store with TRPY1 as Ca2+ release channel (Denis and Cyert, 2002). According to our results TRPY1 currents would be inhibited at 1.3 mM Ca2+, but a considerable amount of Ca2+ is bound by vacuolar polyphosphate (Dunn et al., 1994) and therefore most probably not available for channel inhibition. The remaining vacuolar free [Ca2+] of ~30 μM (Dunn et al., 1994) would allow significant TRPY1 currents (Figure 2). Considering that monovalent cations K+ and Na+ are the major charge carriers used to record TRPY1 currents in the yeast vacuole and in HEK-293 cells, Ca2+ storage and release may not represent major functions of vacuoles and TRPY1 channels. Instead, upon hypertonic shock, which activates TRPY1, cation efflux from the vacuole might provide osmolytes to the cytosol and thereby counteracts water deprivation. The vacuolar Ca2+-dependent inhibition of TRPY1 might protect the yeast cell from cytosolic Ca2+ overload after hypertonic-shock-induced channel activation and increase of the vacuolar [Ca2+] subsequent to loss of vacuolar water.

Limitations of the Study
We studied the Ca2+-dependent modulation of the yeast vacuolar TRP channel and identified two aspartate residues in the outer vestibule as essential for Ca2+ binding and Ca2+-dependent inhibition. The structure of TRPY1 is not yet known, and we aligned the S5, pore linker, and S6 structure of TRPY1 according to the published structure of Xenopus TRPV4. Thus the mechanism we discuss for the Ca2+-dependent inhibition of TRPY1 reflects our experimental data along with a TRPY1 model, obtained due to an alignment to the available structure of another TRP protein. The suggested scenario awaits further support by structural data of TRPY1.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Transparent Methods and six figures and can be found with this article online at https://doi.org/10.1016/j.isci.2018.11.037.

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AUTHOR CONTRIBUTIONS
Conceptualization, A. Beck, M.A., and V.F.; Investigation, M.A., A. Beck, H.W., and A. Belkacemi; Resources, G.S., M.J., and V.F.; Project Administration, A. Beck and V.F.; Supervision, A. Beck, A. Bertl, G.S., and V.F.; Writing – Original Draft, A. Beck; Writing – Review and Editing, A. Beck, G.S., and V.F.; Funding Acquisition, A. Beck, G.S., M.J., and V.F.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Identification of Inhibitory Ca$^{2+}$ Binding Sites in the Upper Vestibule of the Yeast Vacuolar TRP Channel

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Figure S1 Vacuolar Ca$^{2+}$-dependent inhibition of TRPY1 at vacuolar pH 5.5, related to Figure 2

(A) Whole-vacuole currents at -80 and 80 mV extracted from 200 ms ramps (0.5 Hz) spanning from 150 to -150 mV, $V_h$=0 mV, plotted versus time, activated by cytosolic Ca$^{2+}$ (1 mM, indicated by the bar) in the absence (black) and presence (red) of 1 mM Ca$^{2+}$ (vacuolar, applied by the patch pipette) at a vacuolar pH of 5.5. (B) Current-voltage relationships (IVs) of maximum Ca$^{2+}$-induced ($I_{max}$) whole-vacuole currents in A. Currents are shown as means ± S.E.M. and IVs as means with the number of measured vacuoles in brackets.
Figure S2 Effects of divalent cations in the cytosol and in the vacuole on TRPY1 currents, related to Figure 2

Whole-vacuole currents at -80 and 80 mV extracted from 200 ms ramps (0.5 Hz) spanning from 150 to -150 mV, \( V_h = 0 \) mV, plotted versus time, activated by 1 mM cytosolic divalent cations (Ca\(^{2+}\), Ba\(^{2+}\), Mn\(^{2+}\), Sr\(^{2+}\), application indicated by the bar) in the absence of vacuolar Ca\(^{2+}\) (A, left) or activated by 1 mM cytosolic Ca\(^{2+}\) (application indicated by the bar) in the absence (black) and presence of 1 mM vacuolar Ca\(^{2+}\), Ba\(^{2+}\), Mn\(^{2+}\) or Sr\(^{2+}\) (C, left) in yeast YVC1 KO cells expressing the wild-type YVC1 cDNA. (Black traces in A and C are the same). The corresponding current-voltage relationships (IVs) are shown next to the currents in A and C. (B, D) Statistics of the current amplitudes at -80 mV corresponding to the currents shown in A and C. All currents are normalized to the size of the vacuole (pA/pF) and are shown as means ± S.E.M. (A, B, C, D), IVs depict means (A, C). The number in brackets denote the number of measured vacuoles. One-way analysis of variance (ANOVA): not significant (ns), * p<0.05, ** p<0.01, *** p<0.001.
Figure S3 TRPY1 wild-type and mutants in yeast vacuoles, related to Figure 4

Current-voltage relationships (IVs; A-E) and current amplitudes (F, G) correspond to the data shown in Figures 4D-H: Current-voltage relationships (IVs) of the basic currents after break-in (I_{basic}; A, B) and maximum activated whole-vacuole currents (I_{max}; C-E) at 1 mM [Ca^{2+}]_{cyt} in the absence (A, C, E) or presence (B, D, E) of vacuolar Ca^{2+} in yeast YVC1 KO cells expressing YVC1 wild-type or mutant cDNAs (Figure 4D-H). IVs are extracted from 200 ms voltage ramps spanning 150 to -150 mV, V_{h}=0 mV. F and G depict the statistics of the basic (F) and maximal Ca^{2+}-activated (G) currents at -80 mV from the experiments in Figure 4D-H in the absence (left, black) and presence (right, red) of 1 mM vacuolar Ca^{2+}. IVs are normalized to the cell size (pA/pF) and are shown as means; bars depict means ± S.E.M. with number of measured cells in brackets. (H) Statistics of the Ca^{2+} signals measured by relative luminescence units (RLU) in Figure 4I shown as means ± S.E.M. with the number of experiments (x) from y transformations (x/y). One-way analysis of variance (ANOVA): not significant (ns), * p<0.05, ** p<0.01, *** p<0.001 compared to WT.
Figure S4 Vacuolar currents in YVC1 KO yeast expressing wild-type or mutant YVC1 cDNAs (low expression levels) and single channel currents, related to Figure 4

(A-C) Whole-vacuole currents at -80 and 80 mV extracted from 200 ms ramps (0.5 Hz) spanning from 150 to -150 mV, $V_e$=0 mV, plotted vs. time, activated by cytosolic 1 mM Ca$^{2+}$ at 0 (black) or 1 mM (red) vacuolar (patch pipette) Ca$^{2+}$ in yeast YVC1 KO cells expressing TRPY1 wild type (WT; A), D401A (B) or D405A (C) cDNAs. (Plasmids lacking YVC1 promoter and termination sequence => low expression level of YVC1 WT and mutants => low current amplitudes compared to Figure 4). Bars indicate application of 1 mM Ca$^{2+}$. Currents are normalized to the cell size (pA/pF) and are shown as means ± S.E.M.. The number in brackets denote the number of measured cells (x). (D) Representative current traces from voltage steps to 40 mV and -40 mV applied to excised (outside-out) vacuolar patches of yeast expressing TRPY1 WT, TRPY1 D401A and TRPY1 D405A in the absence (left) and presence (right) of 1 mM vacuolar Ca$^{2+}$ (dashed lines = no channels open). (E) Single channel current amplitudes of TRPY1 WT, TRPY1 D401A and TRPY1 D405A in the absence and presence of vacuolar 1 mM Ca$^{2+}$, analyzed at 40 mV from current traces of voltage steps in whole-vacuoles and excised (outside-out) vacuolar patches. Single channels of TRPY1 D405A were analyzed from yeast cells transfected with single copy pRS316 CEN plasmids without its own promoter (see C). Current amplitudes in E are shown as means ± S.E.M. with number of independent experiments indicated. One-way analysis of variance (ANOVA): p<0.05, ** compared to WT in the absence of vacuolar Ca$^{2+}$. 
Figure S5 TRPY1 wild-type and mutants in HEK-293 cells, related to Figure 5

Current-voltage relationships (IVs; A-D and G-I) and current amplitudes (E-F and J) correspond to the data shown in Figures 5B-G and H-J, respectively: Current-voltage relationships (IVs) of the basic current after break-in ($I_{\text{basic}}$; A, B) and maximum 0.75 M sorbitol-induced ($I_{\text{max}}$; C, D) whole-cell currents in the absence (A, C) or presence (B, D) of extracellular 1 mM Ca$^{2+}$ in HEK cells mock-transfected (empty vector; con, control), and transfected with wild-type or mutant YVC1 cDNAs (as in Figures 5B-G). (G-I) Current-voltage relationships of 3 mM cytosolic Ca$^{2+}$-induced whole-cell currents in the absence (black) or presence (red) of extracellular Ca$^{2+}$ in HEK cells transfected with wild-type or mutant YVC1 cDNAs (as in Figures 5H-J). IVs are extracted from 400 ms voltage ramps spanning from -100 to 100 mV, $V_h$=0 mV, normalized to the cell size (pA/pF) and shown as means. E and F depict the statistics of the basic (E) and maximal 0.75 M sorbitol-induced (F) currents at 80 mV from the experiments in Figures 5B-G and J the statistics of the 3 mM cytosolic Ca$^{2+}$-induced current (180 s after break-in) in Figures 5H-J in the absence (left, black) and presence (right, red) of 1 mM (E, F) or 2 mM (J) extracellular Ca$^{2+}$. Bars show means ± S.E.M.. The number in brackets denote the number of measured cells. One-way analysis of variance (ANOVA): not significant (ns), * p<0.05, *** p<0.001 compared to WT (E, F) or compared to 0 ext. Ca$^{2+}$ (J).
Figure S6  S5-S6 linker sequence alignment and structural modelling of TRPY1, related to Figure 6

(A) The S4-S5 linker sequence of *Saccharomyces cerevisiae* TRPY1 (scTRPY1) and the S4-S5 linker sequences defined by the structures of rat TRPV6 (rTRPV6) (Singh et al., 2017), xenopus TRPV4 (xTRPV4) (Deng et al., 2018), rat and human TRPV1 (rTRPV1, hTRPV1) (Liao et al., 2013), rat TRPV3 (rTRPV3), rabbit TRPV2 (rbTRPV2) (Zubcevic et al., 2016) and human TRPA1 (hTRPA1) (Paulsen et al., 2015) were aligned by MUSCLE 3.8. Green boxes mark the selectivity filters. (B, C) Models for TRPY1 based on xenopus TRPV4 ((Deng et al., 2018), PDB: 6BBJ). For the structural alignment, residues 601-700 in TRPV4 were replaced by residues 367-466 from TRPY1. The aspartate residues 398, 401, 405, and glutamate residues 428 and 429 facing the yeast vacuole lumen are indicated by red sticks, Ca^{2+} by red balls. (B) Side view of the S5, pore loop and S6 domains of two opposite TRPY1
wild-type subunits in the absence (left) and presence (right) of Ca\textsuperscript{2+}. (C) Top view of the S5, pore loop and S6 domains of four TRPY1 wild-type subunits in the absence and presence of Ca\textsuperscript{2+} (left), and four non-Ca\textsuperscript{2+} binding TRPY1\textsubscript{D401A} and TRPY1\textsubscript{D405A} subunits (right). Note, that the putative lower gate, represented by the residues I455 and Y458 (see B), narrows significantly upon Ca\textsuperscript{2+} binding at residues D401 and D405 in TRPY1 wild-type (B, C).
Transparent Methods

Yeast cell cultures and expression plasmids

For patch clamp and Ca\(^{2+}\) Imaging, *Saccharomyces cerevisiae* cells (wild-type strain: W303; YVC1-deficient strain: YVC1::TRP1 in W303, GSY1180) were cultured overnight in liquid YPD or SD media (both Sigma Aldrich, St. Louis, US) at 30°C with rotary shaking. The cells were harvested at an OD\(_{600}\) between 1.2 and 1.8. The YVC1 gene encoding wild-type TRPY1 (NCBI accession number NM_001183506.1) was subcloned into the pRS316 vector (Sikorski and Hieter, 1989) with its own promoter and terminator sequence. All expression plasmids used in this study for transformation in YVC1-deficient yeast cells are summarized in Table 1. For Ca\(^{2+}\) imaging, these cells were co-transformed with plasmid pEVP11-AEQ89 (Batiza et al., 1996) encoding the Ca\(^{2+}\) indicator aequorin.

Antibodies, Western blots and cell surface biotinylation

The rat monoclonal anti-TRPY1 antibody (Chang et al., 2010) was generated in-house using amino acid residues 577 to 675 (C-terminus) of the TRPY1 protein, affinity-purified and used at a dilution of 1:500 for Western blotting. Cell lysates were prepared (Wright et al., 1989), proteins separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis and blotted onto polyvinylidene difluoride (PVDF) membranes (Thermo Fisher Scientific, Schwerte, Germany). Proteins were detected with horseradish peroxidase-coupled secondary antibodies and the Western Lightning Chemiluminescence Reagent Plus (Perkin Elmer) for HEK-293 cells and the SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Fisher Scientific) for yeast. Original scans were saved as TIFF files from LAS 3000 (Fujifilm), which were further processed in Adobe Photoshop. Images were cropped,
resized proportionally, and brought to the resolution required for publication. For yeast TRPY1 wild-type and mutant protein densitometric quantification, backgrounds of the Western blots were subtracted and TRPY1 signal intensities were normalized to the signal intensity of the loading control SRP1. The following additional antibodies were used (dilution, company): Anti-rat peroxidase-conjugated goat antibody (1:2,000, Sigma-Aldrich, Taufkirchen, Germany). anti-aequorin rabbit polyclonal antibody (1:1,000, Abcam, Cambridge, UK), anti-rabbit goat peroxidase-conjugated IgG (1:10,000, Santa Cruz Biotechnology, Dallas, TX, USA), anti-Srp1 rabbit antibody (1:1,000, (Gorlich et al., 1996)). Cell-surface protein biotinylation was essentially performed as described in (Fecher-Trost et al., 2013).

Ca²⁺ imaging in yeast cells

The aequorin photoprotein was used to measure cytosolic free [Ca²⁺] in yeast cells (Denis and Cyert, 2002). Cells were resuspended in fresh medium to a density of OD₆₀₀=10. Coelenterazine (Synchem, Felsberg, Germany) was added to a cell suspension of 700 to 1,000 µL at a final concentration of 60 µM. After incubation for 20 min at 30°C, cells were pelleted and resuspended in 700-1,000 µL of fresh medium and incubated again for 45-90 min at 30°C on a roller. 100 µL of the cell suspension were placed into microplate wells and the basal level of luminescence was detected for 30 s at 30°C using a microplate reader (Infinite M200, Tecan, Männedorf, Switzerland). Thereafter 100 µL media containing 3 M NaCl were added yielding a final NaCl concentration of 1.5 M (“hyperosmotic shock”). The luminescence intensity was monitored at 470 nm and plotted as relative luminescence units (RLU) over time using the i-control 1.7 microplate reader software (Tecan). The relative luminescence units obtained for TRPY1 mutants were
normalized to the luminescence units obtained by cells expressing the wild-type YVC1 cDNA at the same day.

*Preparation of giant vacuoles, and whole-vacuole and excised vacuolar patch clamp*

Spheroplasts were prepared as described (Bertl and Slayman, 1990). Briefly, yeast cells were incubated in 3 mL of incubation buffer (50 mM KH₂PO₄, 0.2% β-mercaptoethanol, pH 7.2) for 15 min. In order to remove the cell wall 4 mL of protoplasting buffer (50 mM KH₂PO₄, 0.2% β-mercaptoethanol, 2.4 M sorbitol, pH 7.2) including 150 mg bovine serum albumin (fraction V protease-free; Carl Roth, Karlsruhe, Germany) and zymolyase 20T (ICN Biochemicals, Costa Mesa, USA) to a final concentration of 1 mg/mL were added. After 45 min incubation at 30°C on a roller, spheroplasts were harvested and resuspended in stabilizing buffer (220 mM KCl, 10 mM CaCl₂, 5 mM MgCl₂, 5 mM 2(N-Morpholino)ethanesulfonic acid (MES), 1% (w/v) glucose, pH 7.2). Spheroplasts were incubated for 1 to 2 days to expand and form large vacuoles. The plasma membranes were released by releasing buffer (100 mM potassium citrate, 5 mM MgCl₂, 10 mM glucose, 10 mM MES, pH 6.8). After washing with bath solution (see below), the vacuoles were used for whole-vacuole patch clamp experiments.

Break-in was performed by short voltage pulses (850-1,100 mV, 1-5 ms) and currents were recorded using an EPC-9 patch clamp amplifier (HEKA, Lambrecht, Germany). Experiments were performed at an axiovert 135 microscope (Zeiss, Oberkochen, Germany) equipped with a 40x LD Achroplan objective (Zeiss), a 470 nm LED (Rapp OptoElectronics, Hamburg, Germany) and a GFP filter set (Zeiss). Patch pipettes were pulled from glass capillaries GB150T-8P (Science Products, Hofheim, Germany) with a PC-10 micropipette puller (Narishige, Tokyo, Japan) and
after filling with internal solution had resistances between 2 and 4 MΩ. Both internal (patch pipette = vacuolar) and external (bath = cytosolic) saline contained 150 mM KCl, 5 mM MgCl₂, 2 mM dithiothreitol (DDT), 10 mM HEPES, pH 7.2. In some experiments, various concentrations of CaCl₂, BaCl₂, SrCl₂, or MnCl₂ were added either in the patch pipette, or in an application pipette for direct application onto the measured vacuole. For K⁺-free and low Cl⁻ conditions, KCl was substituted by tetraethylammonium (TEA) chloride or potassium gluconate, respectively.

Voltage ramps of 200 ms spanning from 150 to -150 mV were applied every 2 s from a holding potential (Vₜₐₜ) of 0 mV. From the individual ramp recordings, inward and outward currents were extracted at -80 mV and 80 mV, respectively, and plotted versus time. Representative current-voltage relationships (IVs) were extracted at the stable phase of the current. Currents were normalized to the vacuolar capacitance, which was extracted as a representative measure for the size of the vacuole to calculate current densities (pA/pF). Single channel current amplitudes were measured at -40 mV from current traces of voltage ramps (150 to -150 mV, 200 ms, Vₜₐₜ 0 mV) and voltage steps (-40 mV, 2 s, Vₜₐₜ 0 mV), and at 40 mV from current traces of voltage steps (40 mV, 2 s, Vₜₐₜ 0 mV) in whole-vacuoles and in excised (outside-out) vacuolar patches. Please note that the membrane potentials refer to the cytosolic side, i.e. inward currents at -80 mV represent the movement of positive charges from the vacuole towards the cytosol (Bertl et al., 1992).

**Site-directed mutagenesis, modelling and transfection of HEK-293 cells**

The YVC1 cDNA, encoding wild-type TRPY1 (NM_001183506.1) was used as template for site-directed mutagenesis using the Q5 site-directed mutagenesis kit (New England BioLabs, Ipswich, USA). All DNAs were sequenced on both strands.
Table 1 summarizes all expression plasmids used in this study. Figure S6B and C were prepared using PyMOL Molecular Graphics System (Version 1.5.0.4, Schrödinger, LLC, New York, NY, USA) by Coot (Emsley et al., 2010), based on the coordinates of *xenopus* TRPV4 (PDB: 6BBJ; (Deng et al., 2018)). The full-length of *xenopus* TRPV4 tetrameric structure was used as template. Amino acids 601 to 700 of TRPV4 were replaced by amino acids 367 to 466 (S5, pore linker, S6) from yeast TRPY1 (wild-type, D401A and D405A) in Coot. The structural model of wild-type TRPY1 was calculated in the absence and presence of Ca$^{2+}$ in close proximity to D401 and D405.

HEK-293 cells (ATCC, CRL 1573) were cultured in DMEM (Dulbecco’s modified eagle medium, Thermo Fisher, Waltham, USA), 10% fetal bovine serum (FBS), 1% penicillin/streptomycin). For patch clamp experiments the HEK-293 cells were transfected with 2 µg of the bicistronic pCAGGS-IRES-GFP vector (mock) or plasmids coding for TRPY1 wild-type or the mutants (see Table 1). Fugene HD (Promega, Madison, USA) Lipofectamine 2,000 or 3,000 (Invitrogen, Carlsbad, USA) and 293 Cell Avalanche (EZ Biosystems, College Park, USA) were used as transfection reagents. For whole-cell patch clamp HEK-293 cells were plated on glass cover slips and used for experiments 24-72 h after transfection. The positively transfected HEK-293 cells were identified by their expression of GFP.

**Whole-cell patch clamp experiments in HEK-293 cells**

Whole-cell currents were recorded using the same system and software as described for yeast vacuolar patch clamp. Voltage ramps of 400 ms spanning from -100 to 100 mV were applied every 2 s from a holding potential of 0 mV. Currents were filtered at 2.9 kHz and digitized at 400 ms intervals. From the individual ramp recordings, inward and outward currents were extracted at -80 mV and 80 mV,
respectively, and plotted versus time. Representative current-voltage relationships (IVs) were extracted at the indicated time points. Currents were normalized to the cell capacitance to calculate current densities (pA/pF). Patch pipettes were filled with internal saline comprising 120 mM Cs-glutamate, 8 mM NaCl, 1 mM MgCl₂, 10 mM Cs-BAPTA, 10 mM HEPES (pH 7.2). Ca²⁺ concentrations in the pipette solution were adjusted by the combination of 10 mM Cs-BAPTA with 3.1, 8.2, 9.3, and 9.8 mM CaCl₂ to reach final free Ca²⁺ concentrations of 100 nM, 1 µM, 3 µM, and 10 µM, respectively, calculated by WEBMAXC STANDARD (www.stanford.edu). The external solution comprised 140 mM NaCl, 2.8 mM KCl, 2 mM MgCl₂, 10 mM HEPES, 10 mM glucose (pH 7.2) with or without 1 or 2 mM CaCl₂. Hyperosmotic shock was applied directly onto the measured cells using an application pipette containing 0.5 or 0.75 M sorbitol solved in external solution.

⁴⁵Ca²⁺ binding

25-mer peptides as indicated were synthesized (Intavis ResPepSL peptide spot synthesizer) and spotted at approximately 16 nmoles per spot) onto hardened cellulose membranes. The membranes were activated by methanol (5 min), washed with distilled water and soaked 120 min in buffer containing 60 mM KCl, 5 mM MgCl₂ and 10 mM imidazole-HCl, (pH 6.8), changed every 30 min. Then, membranes were incubated for 30 min in the latter buffer containing additional 1.5 µM (1mCi/L) ⁴⁵Ca²⁺, subsequently rinsed with 50% ethanol for 5 min and dried at room temperature for 3 h. After exposure of the dried membrane to a phosphorimager screen (BAS-IP MS 2040, Fujifilm, Japan) for 12-24 h the screen was scanned by a Typhoon imager (Typhoon FLA 9500, GE Healthcare Life Sciences, USA)

Analysis and Statistics
Patch clamp data were first analyzed in Patchmaster or Fitmaster (HEKA), then transferred to IgorPro (WaveMetrics, Portland, USA) for further analysis and graphical presentation. Yeast luminometric Ca\(^{2+}\) imaging data were transferred as Excel files to IgorPro to prepare graphs. IgorPro or GraphPad PRISM (GraphPad, La Jolla, USA) were used to prepare the bar graphs and to test for statistically significant differences between the means of the independent groups with one-way analysis of variance (ANOVA). ns, *, **, *** represents p values of p > 0.05, p ≤ 0.05, p ≤ 0.01, p ≤ 0.001, respectively. The error bars represent the standard error of the mean (S.E.M.). Final figures were prepared in CorelDRAW (Corel Corporation, Ottawa, Canada).
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