TRPM4 links calcium signaling to membrane potential in pancreatic acinar cells

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Transient receptor potential cation channel subfamily M member 4 (TRPM4) is a Ca2+-activated nonselective cation channel that mediates membrane depolarization. Although, a current with the hallmarks of a TRPM4-mediated current has been previously reported in pancreatic acinar cells (PACs), the role of TRPM4 in the regulation of acinar cell function has not yet been explored. In the present study, we identify this TRPM4 current and describe its role in context of Ca2+ signaling of PACs using pharmacological tools and TRPM4-deficient mice. We found a significant Ca2+-activated cation current in PACs that was sensitive to the TRPM4 inhibitors 9-phenanthrol and 4-chloro-2-[[2-(2-chlorophenoxy)acetyl]amino]benzoic acid (CBA). We demonstrated that the CBA-sensitive current was responsible for a Ca2+-dependent depolarization of PACs from a resting membrane potential of −44.4 ± 2.9 to −27.7 ± 3 mV. Furthermore, we showed that Ca2+ influx was higher in the TRPM4 KO- and CBA-treated PACs than in control cells. As hormone-induced repetitive Ca2+ transients partially rely on Ca2+ influx in PACs, the role of TRPM4 was also assessed on Ca2+ oscillations elicited by physiologically relevant concentrations of the cholecystokinin analog cerulein. These data show that the amplitude of Ca2+ signals was significantly higher in TRPM4 KO than in control PACs. Our results suggest that PACs are depolarized by TRPM4 currents to an extent that results in a significant reduction of the inward driving force for Ca2+. In conclusion, TRPM4 links intracellular Ca2+ signaling to membrane potential as a negative feedback regulator of Ca2+ entry in PACs.

Pancreatic acinar cells (PACs) are the major cell types of the exocrine pancreas. They are responsible for secretion of digestive enzymes and primary fluid. Stimulation by endogenous secretagogues, such as acetylcholine and cholecystokinin, causes inositol 1,4,5-trisphosphate (IP3) generation, and consequent Ca2+ release from the endoplasmic reticulum (ER) through the IP3 receptor (IP3R) Ca2+ channels. The subsequent increase in intracellular Ca2+ concentration ([Ca2+]i) triggers the exocytosis of digestive enzymes (1–4). During this process, termed stimulus–secretion coupling, changing [Ca2+]i may exhibit various spatiotemporal patterns, depending on the magnitude of secretagogue stimulation, which eventually determines the quality and quantity of secretion. Threshold concentrations of secretagogues induce transient and repetitive elevations (oscillations) of [Ca2+]i, highly localized to the apical pole of PAC, which was demonstrated to elicit exocytosis of enzyme containing vesicles (5–9). The spatial limitation of Ca2+ release was explained by the higher density of IP3Rs in this region and the large Ca2+ buffering capacity of a mitochondrial belt surrounding the apical area (10, 11). Higher secretagogue concentrations cause higher [Ca2+]i, that breaks through the mitochondrial firewall and generates propagating Ca2+ waves, which initiate transepithelial fluid secretion as well (12–14). These patterns of Ca2+ signals represent the physiological function of Ca2+ signaling, whereas unduly high concentrations of secretagogues initiate a pathological chain of reactions, beginning with an initial [Ca2+]i peak, followed by a lower, but sustained Ca2+ plateau (9, 15). These, peak plateau-type signals overload the cell with excess amount of Ca2+, which is enough to trigger intra-acinarzymogen activation, self-digestion, leading to acute pancreatitis (16–18). However, both long-lasting oscillatory- and peak–plateau-type Ca2+ signals require Ca2+ influx from the extracellular environment (19–22). The mechanism for Ca2+ entry may be either store independent or store-operated Ca2+ entry (ISOCE) or capacitative Ca2+ entry (23–26). The trigger for SOCE is the significant depletion of the ER Ca2+ content, and its role is to fuel further Ca2+ release during strong stimulation. Otherwise, either type of Ca2+ entry channels are assembled from different isoforms of the ORAI protein, with possible contribution of transient receptor potential canonical 3 (TRPC3) channels to SOCE (27–29).

Following [Ca2+]i elevation, Ca2+ is pumped out from the cytosol by the plasma membrane Ca2+ ATPase (PMCA) or transferred back to the ER by the sarco-ER Ca2+ ATPase (SERCA) (30–33).

Since the spatiotemporal characteristics of Ca2+ signaling fundamentally determine cell behavior and disordered Ca2+...
signaling is directly linked to pancreatic pathology, the major challenge of research in this field is to learn more about the regulation of \([\text{Ca}^{2+}]\), and to find new pharmacological targets and compounds to prevent \(\text{Ca}^{2+}\) overload \((34, 35)\). In our efforts to find new effectors of \(\text{Ca}^{2+}\) signaling in PACs, we examined whether \(\text{Ca}^{2+}\)-regulated ion channels from the transient receptor potential family (transient receptor potential cation channel subfamily M member 4 [TRPM4] and transient receptor potential cation channel subfamily M member 5 [TRPM5]) are expressed in PACs and whether they affect \(\text{Ca}^{2+}\) signaling. First, we performed a comprehensive quantitative PCR (qPCR) analysis using murine pancreas and got a positive result for the TRPM4.

TRPM4 is an \([\text{Ca}^{2+}]\)-activated nonselective cation channel mediating a significant amount of depolarizing current in several cell types \((36)\). Accordingly, plasma membrane depolarization because of TRPM4 activation was demonstrated to control various physiological processes through the activation of voltage-gated \(\text{Ca}^{2+}\) channels (in breath pacemaker neurons and cerebral arterioles) or by decreasing capacitative \(\text{Ca}^{2+}\) entry by limiting the electrochemical driving force for \(\text{Ca}^{2+}\) influx (in mast cells and T-lymphocytes) \((37–40)\). A cation current with the hallmarks of TRPM4 was also reported in PAC by Maruyama and Petersen in 1982 \((41, 42)\); however, studying the role of the current in PAC function was impeded by the lack of pharmacological and genetic tools at that time. Nevertheless, the cation current was suggested to be responsible for the \(\text{Ca}^{2+}\)-dependent transepithelial Na\(^+\) and water transport required for fluid secretion. Importantly, the fact that the current is controlled by extracellular \(\text{Ca}^{2+}\) implies that it may serve as a negative feedback regulator of \(\text{Ca}^{2+}\) influx. Presuming that the inward cation current was carried by TRPM4, its role in the feedback regulation of \(\text{Ca}^{2+}\) influx was tested in this study.

**Results**

In preliminary experiments, RT-qPCR analysis was performed from murine whole pancreas lysates using DNA primer probes designed to recognize the two types of \(\text{Ca}^{2+}\)-dependent cation channels TRPM4 and TRPM5. Parallel experiments were done using primer pairs against the three isoforms of IP\(_3\)R, the three isoforms of the ryanodine receptor (RyR) \(\text{Ca}^{2+}\) release channel and TRPC3, which served as internal control. The housekeeping gene GAPDH expression was used as reference (Fig. 1). IP\(_3\)R isoforms showed high and identical expression, whereas RyR expression was relatively low, with RyR1 being the major isoform. These results are in accordance with previous results showing that all IP\(_3\)R isoforms are equally highly expressed and that RyR has only a complementary role in the \(\text{Ca}^{2+}\) signaling of PACs \((13, 14, 43–45)\). TRPM4 expression was comparable to IP\(_3\)R expression and was significantly higher than that of TRPC3, the \(\text{Ca}^{2+}\) permeable channel partially responsible for SOCE in PACs \((29)\). TRPM5 expression level fell below the detection threshold. As TRPM5 has been shown to be highly expressed in the endocrine pancreas \((46)\), our negative result also implies that mRNA contamination of our whole pancreas lysate by Langerhans islets did not bias our data. Therefore, we conclude that TRPM4 mRNA is highly expressed in PAC.

In the next series of our experiments, functional expression of TRPM4 was tested using the voltage-clamp method in whole-cell configuration of the patch clamp technique. The ionic composition of the extracellular and intracellular solutions was designed specifically for the measurement of nonselective cation currents. To this end, much of \(\text{Cl}^-\) was replaced by glutamate in the recording medium, and Cs\(^+\) was added to the intracellular solution in order to prevent \(\text{Cl}^-\) and \(\text{K}^+\) currents, respectively. Averaged current traces, recorded in course of voltage ramp protocols applied between \(-60\) and \(+120\) mV, are shown in Figure 2, A and B. Under control conditions, a cation current appeared as a small inward background current displaying a slight voltage dependence at positive voltages. When the same cell was treated with cyclopiazonic acid (CPA), a specific inhibitor of SERCA \((47)\), the current significantly increased. CPA is a common tool used to create elevated \([\text{Ca}^{2+}]\), as it inhibits \(\text{Ca}^{2+}\) reuptake and leaves resting ER \(\text{Ca}^{2+}\) leak uncompensated. As a result, CPA treatment raises \([\text{Ca}^{2+}]\) \((35)\). The reason why we used CPA instead of secretagogue stimulation for this purpose is that CPA increases \([\text{Ca}^{2+}]\) while saving the PIP\(_2\) content of the plasma membrane, so it prevents rundown of TRPM4 current during the experiment \((48)\). After reaching steady-state current, cells were treated with a solution containing CPA together with the widely used TRPM4 inhibitor 9-phenanthrol (9-ph; 100 \(\mu\text{M}\)) \((49)\), which diminished the current (Fig. 2A). Unfortunately, 9-ph is not a fully selective inhibitor of TRPM4 as it is known to suppress also the \(\text{Ca}^{2+}\)-dependent \(\text{Cl}^-\) current \((50)\). This issue makes 9-ph problematic to apply with PACs, as both currents show significant depolarizing capacity in these cells. In order to overcome selectivity problems, a more specific TRMP4 inhibitor, 4-chloro-2-[[2-(2-chlorophenoxy)acetyl]amino]benzoic acid (CBA), was applied. About 10 \(\mu\text{M}\) of CBA inhibited
the Ca\(^{2+}\)-dependent cation current as potently as 100 μM 9-ph (Fig. 2B) (49) without affecting the Cl\(^-\) current (Fig. 3, A and B), indicating that CBA is an appropriate drug for selectively inhibiting TRPM4 even under the experimental conditions of live-cell Ca\(^{2+}\) imaging, when both cation and anion currents operate simultaneously in PACs.

Cation current was also measured in PACs isolated from animals in which the gene encoding the TRPM4 was disrupted (TRPM4 KO) (39), using the same experimental arrangement. Although, the mean current was somewhat higher comparing to control, it failed to increase during CPA treatment and was not sensitive to CBA either (Fig. 2C). These results strongly suggest that TRPM4 is functionally expressed in WT PACs in significant amounts.

In order to determine the impact of TRPM4 current on membrane potential, perforated patch clamp experiments were performed using the current clamp technique. The membrane potential was −44.4 ± 2.9 mV under control conditions, and the membrane slowly depolarized to −27.7 ± 3 mV when 30 μM CPA was applied. The membrane potential returned close to the resting value (−42.9 ± 1.6 mV) when the perfusion solution was supplemented with 10 μM CBA (Fig. 4, A and B). Based on these results, we propose that PAC plasma membrane depolarizes in a Ca\(^{2+}\)-dependent manner, involving the activation of TRPM4.

Furthermore, we hypothesized that the driving force for Ca\(^{2+}\) influx at these depolarized membrane potentials is low enough to significantly decrease Ca\(^{2+}\) entry. To test this hypothesis, ratiometric Ca\(^{2+}\) imaging was performed in clusters of PACs, which were exposed to long-term stimulation with low dose (10 pM) of cerulein, which is equivalent to a physiological stimulation with cholecystokinin (51). Parallel experiments were performed in Ca\(^{2+}\)-containing and Ca\(^{2+}\)-free solutions (when the source of Ca\(^{2+}\) can be only intracellular) using control and TRPM4 KO cells as well. Continuous application of 10 pM cerulein evoked periodic fluctuation (oscillation) of [Ca\(^{2+}\)]\(i\), in all groups (Fig. 5, A and B). Ca\(^{2+}\) spikes emerging between 8 and 10 min of cerulein treatment were analyzed because Ca\(^{2+}\) entry was expected to already contribute to the Ca\(^{2+}\) signaling by this time (21). While the average amplitude of Ca\(^{2+}\) spikes in control PACs was very similar in Ca\(^{2+}\)-containing and Ca\(^{2+}\)-free media, the value was markedly higher in Ca\(^{2+}\)-containing medium in the case of TRPM4 KO preparations (Fig. 5C; ΔR\(_{500-600}\) values: CTRL 0 Ca\(^{2+}\): 0.111 ± 0.008; CTRL 2.5 Ca\(^{2+}\): 0.091 ± 0.007; TRPM4 KO 0 Ca\(^{2+}\): 0.087 ± 0.004; and TRPM4 KO 2.5 Ca\(^{2+}\): 0.14 ± 0.011). These data indicate that Ca\(^{2+}\) entry is significant after 8 min cerulein treatment in TRPM4 KO PACs but not in control cells. Importantly, the spike amplitudes in control and TRPM4 KO PACs were essentially the same in Ca\(^{2+}\)-free saline solution, suggesting that the Ca\(^{2+}\) content of ER was similar at the end of cerulein treatment in both types of cells. Area under the curve (which is believed to be proportional to the sum of intracellular Ca\(^{2+}\)) followed a same trend, but the differences were not different significantly (data not shown). Otherwise, the temporal characteristics of Ca\(^{2+}\) transients were not apparently different in control and TRPM4 KO PACs.
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Figure 3. CBA does not affect the Cl− current in pancreatic acinar cells. A, representative current traces of whole-cell currents of a cell under control conditions and during the application of 10 μM CBA. Step depolarizations were applied between −60 and +120 mV with 1 μM Ca2+ in the pipette solution. Averaged data are shown in panel B (n = 5). CBA, 4-chloro-2-[[2-(2-chlorophenoxy)acetyl]amino]benzoic acid.

Figure 4. Ca2+-dependent depolarization of pancreatic acinar cells relies on TRPM4 activity. A, representative membrane potential (MP) recording obtained under current-clamp conditions in control extracellular solution, during CPA (30 μM) and CPA + CBA (10 μM) treatment, respectively. Averaged membrane potentials recorded at different experimental conditions are shown in panel B. Mean values were compared with repeated-measures ANOVA, and pairwise comparisons between the indicated groups were carried out using paired-sample t tests with Bonferroni correction. Asterisks indicate significant (p < 0.05) differences (n = 5). CPA, 4-chloro-2-[[2-(2-chlorophenoxy)acetyl]amino]benzoic acid; CBA, cyclopiazonic acid; TRPM4, transient receptor potential cation channel subfamily M member 4.

Conclusion, the difference between control and KO PAC Ca2+ spikes in Ca2+-containing solution indicates that TRPM4 is likely involved in the negative feedback regulation of Ca2+ entry in PACs. Unfortunately, testing the role of TRPM4 using CBA in a similar experimental setting was not possible because of a strong off-target effect, that is, 30 μM CPA completely abolished Ca2+ oscillations in Ca2+-free bath solution (data not shown), suggesting that CBA inhibited Ca2+ release in PACs.

The hypothesis that TRPM4 is involved in the negative feedback regulation of Ca2+ entry was further investigated in experiments designed to cause significant Ca2+ depletion from the ER in order to turn SOCE on. Therefore, PACs were stimulated with 20 pM cerulein for 10 min in Ca2+-free external solution. Cerulein treatment caused rather sustained Ca2+ signals with fluctuations of gradually decreasing amplitudes (Fig. 6A). This behavior is an obvious sign of ER depletion and Ca2+ unloading because of the activity of PMCA. Afterward, the solution was exchanged to a solution containing 2.5 mM Ca2+, which resulted in a tonic elevation of [Ca2+]i, attributed to the activation of SOCE (Fig. 6A). The amplitude of the SOCE-related fluorescence signal ratio (ΔR) was compared with those measured in TRPM4 KO PACs and found to be significantly higher in KO cells (Fig. 6B; ΔRSOCE values: CTRL [WT]: 0.091 ± 0.005; TRPM4 KO: 0.126 ± 0.004).

To verify these results, SOCE was more specifically investigated, by using CPA in order to cause ER depletion in a receptor-independent manner. This experimental approach avoids possible additional reactions that might interfere with the Ca2+-signaling machinery during secretagogue stimulations. About 30 μM of CPA was applied in Ca2+-free saline to induce Ca2+ leak from the ER (Fig. 7A). In the beginning of treatment, [Ca2+]i increased, which was followed by a slow decrease, indicating that the ER depleted and Ca2+ was eliminated from the intracellular space by PMCA. The amplitude of Ca2+ signals was not significantly different between control, CPA-treated, and KO PACs (0.18 ± 0.02, 0.22 ± 0.01, and 0.21 ± 0.03, respectively). After reaching basal fluorescence values, the Ca2+-free solution was replaced by 2.5 mM Ca2+-containing solution, which resulted in a robust increase in [Ca2+]i (Fig. 7A, left panel). Similar experiments were performed in the presence of CBA or using TRPM4 KO PACs. In these experiments, CBA was appropriate to use for the specific inhibition of TRPM4 because the ER was already depleted and the SERCA pump was inhibited; therefore, CBA could not affect Ca2+ release or the content of the ER. Analysis of Ca2+-influx–related alterations of fluorescence intensity ratios revealed that the slope and amplitude of the change of fluorescence (ΔR) was higher in TRPM4 KO PACs compared with control. In addition, CBA treatment significantly enhanced the rate of rise (but not the amplitude) of the fluorescence signal (Fig. 7, B and C) (slope, CTRL: 0.76 ± 0.04; CBA: 1.06 ± 0.03; KO: 1.88 ± 0.1 AU; ΔRSOCE: CTRL: 0.065 ± 0.004; CBA: 0.071 ± 0.002; KO: 0.103 ± 0.005). These data are

\[ \text{CTRL} \quad +\text{CBA} \]

\[ 0.4 \text{ nA} \]

\[ 0.4 \text{ s} \]

\[ 0.6 \]

\[ 0.2 \]

\[ 0.4 \]

\[ 0.6 \]

\[ 0.0 \]

\[ 0.09 \]

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\[ 0.93 \]

\[ 0.95 \]
in accordance with those presented in Figure 6 and support our hypothesis that TRPM4 is a negative feedback regulator of Ca^{2+} entry in mouse PACs.

**Discussion**

In this study, we provide the first direct evidence that the TRPM4 current depolarizes PACs in a Ca^{2+}-dependent manner and acts as a negative feedback regulator of Ca^{2+} entry under physiological conditions. However, our data may also have pathological implications. Ca^{2+} overload of PACs is believed to be the critical early pathological event, leading to premature intracellular zymogen activation, self-digestion, and eventually, acute pancreatitis (16–18). As SOCE is essential to develop sustained and pathological elevation of [Ca^{2+}]_{i} and ORAI1 inhibitors were reported to mitigate the severity of acute pancreatitis, our data raise the possibility that TRPM4 plays a preventive role in the pathophysiology of Ca^{2+} signaling (34). This hypothesis should be further investigated using animal models of the disease. The translational and therapeutic potential of TRPM4 should be also evaluated.

Similar physiological functions of TRPM4 have been observed in T-lymphocytes and mast cells earlier. TRPM4 silencing transformed Ca^{2+} oscillations to sustained elevations of [Ca^{2+}]_{i} and led to increased interleukin-2 production in Jurkat T cells, which is in accordance with the idea that TRPM4 reduces Ca^{2+} entry by depolarizing the plasma membrane and decreasing the driving force for Ca^{2+} influx (39).

Similarly, our results also imply that TRPM4 current suppresses Ca^{2+} entry by creating a depolarized membrane potential, where the driving force for Ca^{2+} entry is lower;
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Figure 7. TRPM4 activity regulates Ca\textsuperscript{2+} entry evoked by CPA treatment. A, Ca\textsuperscript{2+} signals of control, CBA-treated, and TRPM4 KO acinar cells during 30 μM CPA application in Ca\textsuperscript{2+}-free extracellular solution and following a solution change to 2.5 mM Ca\textsuperscript{2+}. Statistics of the slope and amplitude of the signal (ΔR) recorded in 2.5 mM Ca\textsuperscript{2+} (B and C; n = experiments/cells; nCTRL = 5/35; nCBA: 4/29; and nKO: 5/41). Asterisks indicate significant changes (p < 0.05, one-way ANOVA, Bonferroni post hoc) from control values. CPA, cyclopiazonic acid; TRPM4, transient receptor potential cation channel subfamily M member 4.

however, an alternative mechanism is offered by Park et al. (52). They showed that TRPM4 physically interacts with TRPC3, a Ca\textsuperscript{2+} release–activated Ca\textsuperscript{2+} channel, in human embryonic kidney 293T cells, which results in reduction of channel activity. Since TRPC3 is highly expressed in PACs, this is a possible explanation for our results. In addition, although CBA was expected to block TRPM4, the compound failed to increase the amplitude of SOCE significantly, which might also be explained by the allosteric inhibition of TRPC3 by TRPM4. We presume that the inhibitory interaction between TRPM4 and TRPC3 is not affected by CBA; so TRPC3 is still inhibited by TRPM4 in the presence of CBA, which accounts for the unaltered SOCE amplitude. However, another reason might be that the TRPM4 inhibition is not complete in the applied concentration.

Apparently, the Cl\textsuperscript{−} current (mediated by the recently identified TMEM16a) also acts as a significant depolarizing current in PACs (53–58), which raises the question why acinar cells express functionally redundant ionic currents. The Ca\textsuperscript{2+}-dependent cation current was hypothesized earlier to have an additional function as a Na\textsuperscript{+} uptake channel of the basolateral membrane, which would supply a plausible transcellular Na\textsuperscript{+} transport mechanism with Na\textsuperscript{+}. However, current measurements recorded at the equilibrium potential of Cl\textsuperscript{−} did not show increased cation current when [Ca\textsuperscript{2+}]\textsubscript{i} was elevated in the extreme apical region of the cell but only after [Ca\textsuperscript{2+}]\textsubscript{i} was increased in the whole intracellular space (59). Although in the lack of suitable TRPM4 antibodies for our immunofluorescence studies, we failed to demonstrate TRPM4 expression in PACs, this earlier study strongly suggests that Ca\textsuperscript{2+}-activated cation channels are expressed only in the basal region of the plasma membrane. The results of Kasai and Augustine also imply that the apical membrane does not carry significant cation currents. Consequently, transepithelial fluid secretion is driven by a paracellular (not transcellular) Na\textsuperscript{+} transport, that is, TRPM4 does not participate in the fluid secretion process of PACs. Therefore, we conclude that TRPM4 functions only as a complementary depolarizing current, which is specifically localized in order to negatively regulate Ca\textsuperscript{2+} entry in the vicinity of Ca\textsuperscript{2+} release–activated Ca\textsuperscript{2+} channels.

Experimental procedures

PAC isolation

All experiments complied with the Hungarian Animal Welfare Act and the 2010/63/EU guideline of the European Union and were approved by the Animal Welfare Committee of the University of Debrecen.

Two types of mice were used in this study. The standard strain was C57Bl6, and we also used mice in which the gene encoding the TRPM4 was disrupted (TRPM4 KO). About 3- to 6-month-old mice of both genders were sacrificed by cervical dislocation, and the pancreas was removed immediately. Acinar cells were isolated as described earlier (60). The pancreas was injected with 100 U/ml collagenase P, 0.1 mg/ml trypsin inhibitor, and 2.5 mg/ml bovine serum albumin, dissolved in F12/Dulbecco’s modified Eagle’s medium (DMEM). The tissue was incubated in 5 ml of this solution in a shaking water bath at 37 °C for 25 min while continuously gassed with carbogen. The medium was replaced with fresh medium after 10 min. The tissue was dissociated by trituration performed by 4 to 6 cycles of pipetting through a 10-ml serological pipette, then filtered through a 150 μm mesh. Cells were layered on the top of 2 × 5 ml F12/DMEM, containing 400 mg/ml bovine serum albumin and collected by gentle centrifugation. The pellet was washed in 2 ml DMEM and collected by slow centrifugation. Acinar cell clumps were gently resuspended in DMEM and kept at room temperature until use in Ca\textsuperscript{2+} imaging experiments.

In order to gain single acinar cells for electrophysiological measurements, the resulting acinar cell clumps were subjected to an additional digesting step in Ca\textsuperscript{2+} and Mg\textsuperscript{2+}-free PBS containing 100 U/ml collagenase P for 10 min. Thereafter, cells were dissociated with pipetting using a 5-ml serological pipette.

Intracellular Ca\textsuperscript{2+} imaging

Acinar cell clumps were loaded with 2 μM Fura-8 AM (AAT Bioquest) Ca\textsuperscript{2+} sensitive dye for 30 min at room temperature. Acinar cells were plated on glass-bottom dishes (Bioptechs) and allowed to attach to the bottom. Cells were perfused with Tyrode’s solution containing (in millimolar): 140 NaCl, 5 KCl, 2 MgCl\textsubscript{2}, 2.5 CaCl\textsubscript{2}, and 10 Hepes, and pH 7.38. Some experiments were performed with a Ca\textsuperscript{2+}-free Tyrode’s solution, which contained 0.5 EGTA, but no CaCl\textsubscript{2}. Ratiometric measurement of Fura-8 fluorescence was performed at room temperature using a Zeiss Axiovert 135 microscope equipped.
with a 40× Fluor (1.3 numerical aperture) objective. Fura-8 was excited at 360 and 405 nm at 1 Hz using an light-emitting diode light source (FuraLED; Cairn Research Ltd), and the emitted light was passed through a 520-nm longpass filter and collected using a Qimaging Retiga R3 charge-coupled device camera. The imaging hardware setup was controlled by Microram software (an open source software program) through an interface. Fluorescence values were determined using ImageJ (National Institutes of Health) software with Fiji plugins. Fluorescence ratios of emissions elicited by excitations at 360 and 405 nm were calculated after background subtraction in single cells. Changes of ratios (ΔR) were determined for each cell, averaged, and presented as mean ± SEM.

Electrophysiological recordings

Whole-cell currents were recorded at room temperature using an Axopatch 200B amplifier and a Digidata 1320A digitizer (Molecular Devices) at a 50-kHz sampling rate and filtered online at 5 kHz using a low-pass Bessel filter. Patch pipettes of 5 to 7 MΩ resistance were pulled from borosilicate glass capillaries (Warner Instruments). In TRPM4 current measurements, pipettes were filled with a solution containing (in millimolar): 144 Cs-glutamate, 1 MgCl₂, 0.1 EGTA, 0.0486 CaCl₂ (100 nM ionized Ca²⁺), 3 K-ATP, 10 Hepes, at pH 7.3. The external solution contained (in millimolar): 140 sodium glutamate, 4 CsCl, 2 MgCl₂, 10 Hepes, at pH 7.4.

In Cl⁻ current measurements, the pipette solution contained (in millimolar): 140 N-methyl-D-glucamine chloride, 1 MgCl₂, 1.72 CaCl₂, 2 EGTA (1 μM ionized Ca²⁺), at pH 7.2. The extracellular solution contained (in millimolar): 140 N-methyl-D-glucamine chloride, 1 MgCl₂, 5 glucose, 10 Hepes, at pH 7.3. All ingredients of the solutions were purchased from Sigma–Aldrich (Merck).

TRPM4 current was measured using a ramp voltage protocol applied from −100 to +120 mV, whereas Cl⁻ current was recorded during 1 s long step depolarizations applied between −60 and +120 mV. At least 70% of the series resistance was compensated in these measurements.

Membrane potential was measured under current-clamp condition using the perforated patch clamp technique. The cells were bathed in Tyrode’s solution, whereas the tip of the pipette was filled with a solution containing (in millimolar): 85 potassium glutamate, 45 KCl, 15 NaCl, 2 MgCl₂, 0.1 EGTA, 0.0486 CaCl₂ (100 nM ionized Ca²⁺), 10 Hepes at pH 7.3. The pipette was back-filled with the same solution, supplemented with 300 μg/ml amphotericin B.

RNA isolation, RT, and quantitative real-time PCR

qPCR was performed on a Roche LightCycler 480 System (Roche) using the 5′ nuclease assay (63). Total RNA was isolated using TRIzol (Life Technologies Hungary Ltd), DNase treatment was performed according to the manufacturer’s protocol, and then 1 μg of total RNA was reverse-transcribed into complementary DNA using High-Capacity cDNA Kit from Life Technologies Hungary Ltd. PCR amplification was performed using the TaqMan Gene Expression Assays (assay IDs: Mm01175211_m1 for RYR1, Mm00465877_m1 for RYR2, Mm01328421_m1 for RYR3, Mm00439907_m1 for inositol 1,4,5-trisphosphate receptor (ITPR) type 1, Mm00444937_m1 for ITPR type 2, Mm01306070_m1 for ITPR type 3, Mm00613173_m1 for TRPM4, Mm01129032_m1 for TRPM5, and Mm00444690_m1 for TRPC3) and the TaqMan universal PCR master mix protocol (Applied Biosystems). As internal control, transcripts of the housekeeping gene (GAPDH; assay ID: Mm99999915_g1) were determined. The amount of the transcripts was normalized to the housekeeping gene using the ΔCT method.

Chemicals

Fura-8 AM was purchased from AAT Bioquest. CBA was from Tocris Bioscience (Bio-Techne Corporation). All other chemicals (including collagenase P, 9-ph, CPA, and cerulein) were obtained from Sigma–Aldrich (Merck).

Statistical analysis

Analysis was made in Origin 7.0 (Microcal Software) or in Microsoft Excel. Data are presented as the average of cells obtained from at least three independent experiments and at least three animals. Averages are expressed as mean ± SEM. Statistical analysis was performed by using Student’s t test or one-way ANOVA with Bonferroni post-test. Related samples were analyzed using repeated-measures ANOVA, and pairwise comparisons were carried out with Bonferroni-corrected paired sample t test. Differences were considered significant when p was less than 0.05.

The number of experiments (n) denotes the number of experimental repeats/total number of cells in the case of Ca²⁺ imaging and the number of cells in patch clamp measurements as indicated in the legends to the figures.

Data availability

All data are contained within the article and available upon request.

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Abbreviations—The abbreviations used are: 9-ph, 9-phenanthrol; [Ca²⁺], intracellular Ca²⁺ concentration; CBA, 4-chloro-2-([2-(2-chlorophenoxy)acetyl]aminobenzoic acid; CPA, cyclopiazonic acid; DME, Dulbecco’s modified Eagle’s medium; ER, endoplasmic reticulum; IP₃, inositol 1,4,5-trisphosphate; IP₃R, IP₃ receptor; ITPR, inositol 1,4,5-trisphosphate receptor; PAC, pancreatic acinar cell; PMCA, plasma membrane Ca²⁺ ATPase; qPCR, quantitative PCR; RyR, ryanodine receptor; SERCA, sarco-ER Ca²⁺ ATPase; SOCE, store-operated Ca²⁺ entry; TRPC3, transient receptor potential canonical 3; TRPM4, transient receptor potential cation channel subfamily M member 4; TRPM5, transient receptor potential cation channel subfamily M member 5.

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