Atypical properties of a conventional calcium channel β subunit from the platyhelminth Schistosoma mansoni

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Abstract

Background: The function of voltage-gated calcium (CaV) channels greatly depends on coupling to cytoplasmic accessory β subunits, which not only promote surface expression, but also modulate gating and kinetic properties of the α1 subunit. Schistosomes, parasitic platyhelminths that cause schistosomiasis, express two β subunit subtypes: a structurally conventional β subunit and a variant β subunit with unusual functional properties. We have previously characterized the functional properties of the variant CaVβ subunit. Here, we focus on the modulatory phenotype of the conventional CaVβ subunit (SmCaVβ) using the human CaV2.3 channel as the substrate for SmCaVβ and the whole-cell patch-clamp technique.

Results: The conventional Schistosoma mansoni CaVβ subunit markedly increases CaV2.3 currents, slows macroscopic inactivation and shifts steady state inactivation in the hyperpolarizing direction. However, currents produced by CaV2.3 in the presence of SmCaVβ run-down to approximately 75% of their initial amplitudes within two minutes of establishing the whole-cell configuration. This suppressive effect was independent of Ca2+, but dependent on intracellular Mg2+-ATP. Additional experiments revealed that SmCaVβ lends the CaV2.3/SmCaVβ complex sensitivity to Na+ ions. A mutant version of the CaVβ subunit lacking the first forty-six amino acids, including a string of twenty-two acidic residues, no longer conferred sensitivity to intracellular Mg2+-ATP and Na+ ions, while continuing to show wild type modulation of current amplitude and inactivation of CaV2.3.

Conclusion: The data presented in this article provide insights into novel mechanisms employed by platyhelminth CaVβ subunits to modulate voltage-gated Ca2+ currents that indicate interactions between the Ca2+ channel complex and chelated forms of ATP as well as Na+ ions. These results have potentially important implications for understanding previously unknown mechanisms by which platyhelminths and perhaps other organisms modulate Ca2+ currents in excitable cells.

Background

Voltage-gated calcium (CaV) channels couple membrane depolarisation to the entry of Ca2+ that, in turn, is fundamental in a variety of cellular events such as contraction [1,2], changes in gene expression [3] and neurotransmitter release [4,5]. CaV channels belong to the super-family
of voltage-gated ion channels that also include sodium channels and potassium channels [6], and can be broadly classified into high-voltage activated (HVA) and low-voltage activated (LVA) classes. HVA Ca$_2^+$ channels are heteromeric protein complexes composed of a pore-forming $\alpha_1$ subunit and auxiliary $\beta$ and $\alpha_2\delta$ subunits [7]. In addition to promoting surface expression of the Ca$_2^+$,$\alpha_1$ subunit, Ca$_2^+$,$\beta$ subunits modulate the kinetics of activation and inactivation, gating [8-10] and the rate of recovery from inactivation [11,12].

Schistosomes are parasitic trematode flatworms that cause schistosomiasis, a tropical disease affecting approximately 200 million people worldwide. With the ultimate goal of understanding the molecular basis for neuromuscular transmission in these parasitic flatworms, we have previously cloned three transcripts from *Schistosoma mansoni* that code for one L-type-like and two non L-type high voltage-activated Ca$_2^+$ channel subunits [13]. Heterologous expression of these $\alpha_1$ subunits in *Xenopus* oocytes and mammalian cell lines has proved problematic, perhaps because of the relatively high A-T content of these coding regions, or the lack of a specific chaperone in these systems. Additionally, we have identified two Ca$_2^+$ channel $\beta$ subunits from schistosomes and other platyhelminths: a conventional $\beta$ subunit (SmCa$\beta$), and a variant $\beta$ subunit (SmCa$\beta_{var}$), which appears to be unique to platyhelminths and has unusual structural and functional features [14,15].

When heterologously expressed in *Xenopus* oocytes, the conventional schistosome Ca$_2^+$,$\beta$ subunit significantly increases Ca$_2^+$,2.3 current amplitude, and shifts the steady state inactivation curve to more hyperpolarized potentials [16] (in these experiments, we use the robustly expressing human Ca$_2^+$,2.3d $\alpha_1$ subunit as a "reporter" to assess $\beta$ subunit function). The actions of this $\beta$ subunit are consistent with those of mammalian Ca$_2^+$,$\beta$ subunits [10]. Here, we have reproduced and extended our previous data on modulation of Ca$_2^+$,2.3 currents by the schistosome SmCa$_\beta$ subunit (SmCa$\beta$) in a mammalian cell system, which may better approximate the cellular milieu in which these channels are found in situ. *Xenopus* oocytes are widely used in expression of ion channels and other proteins, in part because they are primed for high levels of protein translation. However, most other cells are not that strongly geared towards this role. Oocytes are much larger than adult, differentiated cells and contain high amounts of yolk granules. In addition, the mammalian cell line HEK does not express the endogenous Ca$_2^+$,$\beta$ subunit that complicates analysis of heterologously expressed Ca$_2^+$ channels in *Xenopus* oocytes [17]. Finally, adult *S. mansoni* live in a mammalian host environment.

Interestingly, during these studies, we observed a rapid run-down of the currents produced by Ca$_2^+$,2.3 channels co-expressed with this schistosome Ca$_2^+$,$\beta$ subunit. Decrease of Ca$^{2+}$ channel activity under whole-cell patchclamp, a configuration of the patch-clamp technique that disrupts the contact between membrane and cytoplasm, is a well-known phenomenon [18-21]. However, very few studies have dealt with the structural and biochemical causes for run-down, and those studies focus primarily on L-type Ca$^{2+}$ channels. Notably, Kameyama and collaborators were able to relate run-down of L-type Ca$^{2+}$ channels to the Ca$^{2+}$-binding protein calmodulin [22]. Here we investigate the mechanism of this $\beta$ subunit-dependent rundown, examining the role of several forms of ATP and cations in run-down of the Ca$_2^+$,2.3/SmCa$_\beta$ complex. Additionally, we use a truncated $\beta$ subunit protein to provide clues regarding the molecular substrate within the SmCa$_\beta$ subunit that mediates run-down of Ca$_2^+$,2.3/SmCa$_\beta$ currents.

**Results**

**SmCa$_\beta$ modulates activation and inactivation of Ca$_2^+$,2.3 in a conventional manner**

To assess the modulatory phenotype of Ca$_2^+$,$\beta$ subunits, a HEK-293 cell line stably expressing the human Ca$_2^+$,2.3d subunit (GB Acc. # L27745) was used. Using Ca$^{2+}$ as the charge carrier, currents produced by Ca$_2^+$,2.3 alone peaked at +30 mV with an average amplitude of -261 ± 20 pA (n = 15). Co-expression of SmCa$_\beta$ increased peak amplitudes to -1640 ± 276 pA (n = 5) and shifted the I–V peak leftwards, to +20 mV (Figure 1A,B).

Using Ca$^{2+}$ as the charge carrier, the decaying phase of Ca$_2^+$,2.3 currents produced by Ca$_2^+$,2.3 channels alone or with SmCa$_\beta$ subunits was well fitted by a double exponential function with fast (τ$_{fast}$) and slow (τ$_{slow}$) time constants of inactivation (Table 1). The double exponential fit to the decay of currents produced by Ca$_2^+$,2.3 in response to a pulse to +30 mV had a fast time constant of 16 ms that contributed 66% to total current decay, and a slow time constant of 69 ms that represented 34% of total current decay. The double exponential fit to the decay of currents produced by Ca$_2^+$,2.3 co-expressed with SmCa$_\beta$ in response to a depolarizing pulse to +20 mV had a fast time constant of 31 ms, which contributed 55% to total current decay, and a slow time constant of 116 ms that contributed 45% to total inactivation. Fundamentally, SmCa$_\beta$ slowed macroscopic inactivation of Ca$_2^+$,2.3 currents by increasing the time constants of both the fast and slow components while simultaneously decreasing the contribution of the fast component and increasing the contribution of the slow component to total inactivation (Figure 1C and Table 1). Co-expression of SmCa$_\beta$ markedly shifted the midpoint of steady-state inactivation in the hyperpolarizing direction (Figure 1D). Midpoints of single order Boltzmann fits to steady-state inactivation curves of Ca$_2^+$,2.3 and Ca$_2^+$,2.3 + SmCa$_\beta$ currents were, respectively, -8 mV and -49 mV (P < 0.05, Student t-test). The
**Figure 1**

**SmCavβ modulates Cav2.3 currents in HEK cells.** A. Representative Ca\(^{2+}\) currents generated by voltage steps to -5, +15, +35 and +55 mV from a holding potential of -80 mV in HEK cells expressing Cav2.3 alone and in combination with SmCavβ or Cavβ2a. B. Current-voltage relationships from HEK cells expressing Cav2.3 alone (filled circles) and in combination with SmCavβ (open circles) or Cavβ2a (filled triangles). Peak currents generated by voltage steps from -20 mV to +75 mV in 5 mV steps from a holding potential of -80 mV were measured. Data points represent mean ± s.e.m. N = 5–15. C. Two-variable plot showing the relationships between inactivation time constants and percentages of contribution to total inactivation for Cav2.3 alone (filled circles) and in combination with SmCavβ (open circles). Data points represent mean ± s.e.m. N = 7–9. Slow time constants for the kinetic of recovery are grouped (dotted lines). Fast time constants of recovery from inactivation kinetics are also grouped (dotted lines). D. Voltage-dependence of inactivation (steady-state inactivation) for Cav2.3 alone (filled circles) and in combination with SmCavβ (open circles) or Cavβ2a (filled triangles). Lines represent fits to the Boltzmann function (see Materials and Methods); \(V_{50}\) and slope values are shown in Table 1. Data points represent mean ± s.e.m. N = 3–6. E. Recovery from inactivation from -100 mV for Cav2.3 channels alone (filled circles) or in the presence of SmCavβ (open circles), using Ba\(^{2+}\) as the charge carrier. Solid curves represent single exponential fits to the data. Values are means ± s.e.m. N = 3.
down, indicating that this phenomenon is Ca2+ independent and shifted the I–V peak in the hyperpolarizing direction (Figure 1).

Macroscopic inactivation

|                | Ca2.3 alone | Ca2.3 + SmCaβ |
|----------------|-------------|---------------|
| V50 (mV)       | -8 ± 4 (3)  | -49 ± 3 (3)*  |
| Slope (mV/e-fold) | 12 ± 2 (3)  | 10 ± 1 (3)    |

Recovery from inactivation

|                | Ca2.3 alone | Ca2.3 + SmCaβ |
|----------------|-------------|---------------|
| τ rec-80 mV (ms) | 178 ± 28 (3) | 252 ± 17 (4) |
| τ rec-100 mV (ms) | 71 ± 7 (3)  | 168 ± 16 (3)* |
| τ rec-120 mV (ms) | 38 ± 1 (3)  | 109 ± 10 (3)* |

Data represent mean ± s.e.m. Number of repeats is indicated in parentheses. Asterisks indicate statistically significant difference (Student t test, P < 0.05).

slope factors of these Boltzmann fits to steady-state inactivation were not significantly different: 12 mV/e-fold for Ca2.3 and 10 mV/e-fold for Ca2.3 + SmCaβ (Table 1).

SmCaβ significantly slowed the rate of recovery from inactivation. The fractional recoveries of current as a function of time at -80, -100 and -120 mV were well fitted by a single exponential function with recovery time constants from inactivation (τrec). For example, τrec for Ca2.3 and Ca2.3 + SmCaβ from -100 mV were, respectively, 71 and 168 ms (p < 0.05, Student t-test) (Figure 1E; see Table 1 for recovery time constants for the three potentials).

The mammalian (rat) β2a subunit (GB Acc. # M80545) increased current amplitude to a similar extent as the schistosome Caβ subunit, shifted the steady-state inactivation curve in the hyperpolarizing direction, although to a lesser degree than the schistosome Caβ subunit, slowed macroscopic inactivation, and shifted the I–V peak in the hyperpolarizing direction (Figure 1).

### Ca2.3 currents run-down in the presence of SmCaβ in a manner independent of Ca2+ but dependent on chelated forms of ATP and free Na+

Currents produced by Ca2.3 progressively decrease in amplitude to approximately 75% of their initial values within ~2.5 minutes of establishing the whole-cell configuration, but only when SmCaβ is co-expressed (Figure 2A). Over the same time frame, no run-down was observed for currents produced by Ca2.3 expressed alone, nor in the presence of the mammalian Caβ2a subunit nor with the structurally atypical schistosome SmCaβvar subunit (Figure 2B). Substitution of CaCl2 by an equimolar concentration of BaCl2 did not prevent run-down, indicating that this phenomenon is Ca2+ independent (Figure 2A).

We hypothesized that this rapid run-down could be caused by physical dissociation between SmCaβ and Ca2.3. However, as the kinetics of inactivation and the steady state inactivation properties were identical before and after run-down (Figure 2C,D), it seems likely that the association between Ca2.3 and SmCaβ remained intact in these conditions.

Since Mg2+ is known to block Ca2+ channels from the intracellular side, we replaced Mg2+-ATP with an equimolar concentration (5 mM) of Na2+-ATP, yet under these conditions we still observed significant run-down that, again, was dependent upon co-expression of the schistosome Caβ subunit (Figure 3A). No run-down was observed in whole-cell patch-clamp experiments using a "minimal" internal solution without ATP (Figure 3C). Further experiments indicate that non-chelated forms of ATP (Tris2-ATP) do not mediate SmCaβ-dependent Ca2.3 run-down (Figure 3B). Currents produced by Ca2.3/Caβ2a did not run-down in the presence of 5 mM intracellular NaCl, Na2-ATP, Tris2-ATP or in minimal intracellular solution (Figures 3D1 and 3D2).

Next we explored the possibility that this particular form of run-down was caused by the cations associated with the ATP molecule (Mg2+ and Na+). Free Mg2+ reduced currents produced by Ca2.3 alone by about 75%, and by about 90% when SmCaβ was co-expressed (Figure 4); therefore, since it would be very challenging to distinguish Ca2+ current decrease due to run-down from Ca2+ current decrease due to blockade by internal Mg2+, we did not pursue run-down studies in the presence of free Mg2+. However, in contrast with Mg2+, free Na+ did not suppress the activity of Ca2.3 channels expressed alone or co-expressed with the schistosome SmCaβ or with the mammalian Caβ2a subunit (Figure 5A–D). However, substituting Mg2+-ATP with NaCl resulted in significant run-down (Figure 5E), indicating Na+-dependent modulation of channels containing SmCaβ.

### The structural basis for run-down mediated by SmCaβ resides in its acidic N-terminal domain

The N-terminus of the SmCaβ subunit contains an atypical domain that is rich in glutamic acid and aspartic acid residues (Figure 6A). Since BLAST screens for similar domains in other proteins did not yield any hits, we set to investigate whether this N-terminal domain represents the structural base for the atypical modulation of the SmCaβ subunit. Furthermore, there is precedence for important modulatory effects on Caβ channels localizing to the N-terminal regions of β subunits [23-26]. To this end, we created a truncated version of SmCaβ that lacks the first forty-six amino acids containing this acid-rich domain (Figure 6A), and tested whether it still caused Ca2.3 currents to run-down. The N-terminally truncated SmCaβ retain...
SmCavβ subunit enhanced Ca v2.3 currents, slowed their macroscopic inactivation and shifted their steady-state inactivation to the same extent as the wild type SmCavβ subunit (Figure 6B,C). However, unlike the wild-type version of SmCavβ, the N-terminally truncated SmCa vβ subunit did not induce run-down of Ca2+ currents within 2–3 minutes of establishing the whole-cell configuration with an internal solution containing either 5 mM Mg2+-ATP (Figure 6D,F), or 5 mM NaCl (Figure 6E,F).

Discussion

In this study we set out to characterize the modulatory phenotype of a Ca vβ subunit from the parasitic trematode S. mansoni. This β subunit is phylogenetically close to other Ca vβ subunits [14]. The mammalian HEK-293 cell line was used as the expression system and the mammalian Ca v2.3 channel splice variant “d” as our “test” α1 subunit. This splice variant is susceptible to modulation by Ca vβ subunits [27] and is the longest of all known versions of the Ca v2.3 subunit [28]. Our data show that SmCavβ exhibits similar modulatory phenotypes in HEK cells and Xenopus oocytes. Additionally, SmCavβ slows recovery from inactivation of Ca v2.3 currents using Ba2+ as the charge carrier, i.e. in the absence of intracellular Ca2+ accumulation. Using strontium as the charge carrier, mammalian and jellyfish Ca vβ subunits also slow recovery from inactivation of this same channel in conditions in which Ca2+ does not accumulate in the intracellular

Figure 2

SmCa vβ-dependent run-down of Ca v2.3 currents. A. Relative peak amplitude of currents produced by Ca v2.3 channels alone (circles) and in combination with SmCavβ (triangles), plotted as a function of time. Currents were normalized with respect to those measured immediately after rupturing the patch membrane (t = 0). Ca2+ currents (filled symbols) and Ba2+ currents (open symbols) are shown. Data points represent mean ± s.e.m. B. Relative amplitude of currents produced by Ca v2.3 in combination with Ca vβ2a (squares) or SmCa vβvar (triangles), plotted as a function of time. As in A, currents were normalized with respect of those at time 0, measured immediately after break-in. Data points represent mean ± s.e.m, N = 3 – 5. C. Ca2+ currents generated by Ca v2.3 in the presence of SmCa vβ at t = 0 min and at t = 2 min. Note that the kinetics of macroscopic inactivation remains unchanged. D. Voltage-dependence of inactivation (steady-state inactivation) at t = 0 min (filled circles) and at t = 2 min (open circles). Solid lines represent fits to the Boltzmann function.

Figure 3

Run-down of Ca v2.3/SmCavβ currents in the presence of intracellular forms of ATP. Relative current amplitude of currents produced by Ca v2.3 (filled circles) channels alone and in combination with SmCavβ (open circles) plotted as a function of time in the presence of 5 mM Na2+-ATP in the patch-pipette solution (A), 5 mM Tris2-ATP (B), and using a “minimal” pipette solution, with no ATP (C). Currents were normalized with respect to those measured immediately after rupturing the patch membrane (t = 0). Panels D1 and D2 show current amplitude produced by Ca v2.3/SmCavβ2a channels as a function of time for all conditions explained above. Symbols are as shown on the figure. Values are means ± s.e.m, N = 3 – 8.
compartment [11]. However, it is important to note that recovery from inactivation appears to depend heavily on the identity of the Ca_α_1 subunit being assayed: all jellyfish and mammalian Ca_β subunits delayed recovery from inactivation of Ca_{2,3} channels, but accelerated the recovery of the jellyfish L-type Ca_α channel [11].

**SmCa_β-dependent run-down of Ca_{2,3} currents**

While characterizing the modulatory phenotype of SmCa_β, we consistently observed a rapid run-down of Ca_{2,3} currents only in the presence of SmCa_β. This contrasts with previous studies, in which run-down appears to occur in the Ca_α_1 subunit, independently of accessory subunits [29-31]. Down-regulation of Ca^{2+} currents caused by interaction of Ca_β subunits with intracellular proteins is well documented. For example, interaction of Ca_β subunits with small GTPases of the RGS family [32], with large GTPases of the dynamin family [33], or with the nuclear protein HP1 [34] results in down-regulation of Ca^{2+} currents. Our data represent the first case, to our knowledge, of down-regulation of Ca^{2+} currents by interactions between Ca_β subunits and chelated forms of ATP or Na^{+} ions, which are likely relevant to platyhelminth physiology. Unlike previous studies, in which hydrolysable forms of ATP suppress Ca^{2+} current run-down [35,36], here we show that two physiologically relevant, hydrolysable forms of ATP induce SmCa_β-mediated decrease in Ca_{2,3} activity. The possibility that run-down was caused by Mg^{2+} ions that dissociate from the ATP molecule was considered, but the fact that free Mg^{2+} dramatically suppressed Ca_{2,3} currents posed a significant challenge to explore this possibility. These results are reminiscent of the effects of Mg^{2+} ions at a similar concentration on L-type Ca_α channels, which occurs by their binding to a low-affinity site at the pore of the Ca_α_1 subunit [37]. Since ATP associates with Na^{+} ions under physiological conditions [38], we also measured run-down of Ca_{2,3}/SmCa_β channels in the presence of intracellular Na^{+}-ATP. Because we detected significant run-down under these conditions, and knowing that the binding constant between Na^{+} ions and ATP is relatively low, about 13 M^{-1} (in contrast to 9000 M^{-1} for Mg^{2+}, [39]), we hypothesized that Na^{+} ions that had dissociated from ATP were causing this run-down. Experiments using NaCl instead of Na^{+}-ATP confirmed this hypothesis.

Our study is not the first to show Ca_β-dependent run-down of non-L type Ca^{2+} current. The mammalian Ca_β subunit appears to enhance run-down of the non-L-type, Ca_{2,1} α_1 subunit [40]. However, it is important to note that in that study, run-down occurred under the two-electrode voltage clamp configuration on Xenopus oocytes, where the connection between cytoplasm and plasma membrane remains largely intact. Therefore the mechanism(s) of run-down employed by the mammalian Ca_β subunit is likely to be distinct from that used by the schistosome Ca_β subunit.

**Molecular substrate for SmCa_β-dependent Mg^{2+}-ATP and Na^{+} sensitivity**

Since the conventional SmCa_β subunit contains a string of acidic residues in the N-terminal domain that is not found in its mammalian counterparts, we generated a version of this subunit that lacked this region, to test whether this unique domain is the molecular substrate for this particular form of run-down. Deletion of this acidic, N-terminal fragment did not change the modulatory phenotype of SmCa_β on current amplitude, inactivation kinetics or steady-state inactivation. However, in contrast to our results with wild type SmCa_β, Ca_{2,3} currents did not run down when co-expressed with this mutated subunit, using Ca^{2+} as the charge carrier and in the presence of intracellular free Na^{+} or Mg^{2+}-ATP. It seems likely that sensitivity to intracellular Mg^{2+}-ATP and free Na^{+} resides in all or part of the string of acidic residues of the SmCa_β subunit.

**Physiological relevance**

Several reports have shown that the Ca^{2+} currents of platyhelminth muscle and nerve cells are very labile, running down within minutes or even seconds after establishing the whole-cell patch-clamp configuration. In previous work, the intracellular solution used to record voltage-gated Ca^{2+} currents from muscle cells of *S. mansoni* was titrated with NaOH to bring the pH to a physiological value [41]. We have empirically calculated that this action would bring the concentration of Na^{+} to approximately 19 mM, which is sufficient to reduce Ca^{2+} currents modulated by SmCa_β significantly, according to our data. Similarly, Ca^{2+} currents recorded from isolated muscle cells of the free-living flatworm *Bidelloura candida* run-down within 20 seconds of establishing the whole-cell configuration [42]: It is tempting to speculate that this run-down was caused by the relatively high concentration of NaCl (30 mM) added to the intracellular solution in these experiments.

Our previous studies have identified a different schistosome Ca_β subunit (SmCa_β_{ar} [14]), which does not exhibit the hallmark action of Ca_β subunits, namely, to increase Ca^{2+} current density. Together with the data presented here, this raises the question of whether schistosomes employ unique strategies to modulate excitability via atypical modulation of HVA Ca_α channels, information that could be useful in the design of targeted therapies to treat schistosomiasis.

**Conclusion**

We have identified novel functions for a schistosome Ca_β subunit, namely to confer Ca^{2+} currents with sensitivity to intracellular Mg^{2+}-ATP and Na^{+} ions, which translates into a reduced ability on the part of this Ca_β subunit to
increase Ca\(^{2+}\) currents. We conclude that the molecular basis for this atypical sensitivity to both Mg\(^{2+}\)-ATP and Na\(^{+}\) ions resides in a domain or domains located within the first forty-six amino acids of SmCa\(_{\beta}\), which contains a string of twenty-two aspartic and glutamic acid residues not present in other Ca\(_{\beta}\) subunits.

**Methods**

**Materials**

Tissue culture dishes were purchased from Corning (NY, USA), Dulbecco's modified Eagle's media (DMEM) was purchased from Invitrogen, poly-L-lysine and ATP salts were purchased from Sigma. The transfection reagent, Tfx, was purchased from Promega. Restriction enzymes were from NEB, and oligonucleotide primers were from MWG Biotech.

**Preparation of eukaryotic expression plasmids encoding SmCa\(_{\beta}\)**

Using standard methods, we cloned all Ca\(_{\alpha}\)\(_{\beta}\) subunits into the pXOOM vector [43], which is optimized for expression of inserts in mammalian cells (under control of a cytomegalovirus promoter), and contains the gene for green fluorescent protein (GFP) as a marker for transfection. For SmCa\(_{\beta}\), the insert from the original SmCa\(_{\beta}\) clone in pCR4-TOPO (Invitrogen) was amplified using Phusion high-fidelity DNA polymerase (NEB). Primers were designed against the beginning and end of the coding regions of the sequence, and included appropriate restriction sites for insertion into pXOOM. The primers were: Forward: 5'-GGGACCTATGCGTGGTACGGAGGATC-3', which includes two G residues and a *Hind III* site at the 5' end; and Reverse: 5'-GGGGCGCGCCGT-TAAATCTGATGACCTGAGCA-3', which includes two G residues and a *Not I* site at the 5' end. Following an initial 98° denaturation for 30 seconds and 25 cycles of 98° for 10 s, 68° for 30 s, and 72° for 2 min, the reaction was purified over a Qiagen spin column (Qiagen), and digested with *Hind III* and *Not I*. The digested band was gel-purified using Quantum Prep Freeze-n-Squeeze columns (BioRad), Pellet Paint (Novagen) was added as carrier, and the product was ethanol-precipitated and ligated to pXOOM that had been digested with *Hind III* and *Not I* and gel-purified. All constructs were sequenced to verify the absence of PCR errors.

**Cell culture and transfection of HEK293-Ca\(_{2.3}\) cells**

HEK293 cells stably transfected with Ca\(_{2.3d}\) [44] were cultured in DMEM supplemented with L-glutamine, glucose and 10% foetal bovine serum in a humidified atmosphere (95%) at 5% CO\(_2\) and 37°C. Cells were used for up to 20 passages and were split every 2–4 days. For electrophysiological recordings, cells were seeded in Petri dishes coated with poly-l-lysine, and transfection of auxiliary \(\beta\) subunits was performed with Tfx on cells at a confluence of 50–60%, using 1 µg of the construct and a DNA: Tfx ratio of 1:2. Cells exhibiting green fluorescence were used for further study.

**Construction of a N-terminal deletion mutant of the SmCa\(_{\beta}\) subunit**

To generate a mutant subunit lacking amino acids 2–46 of the N terminus domain, a diluted sample of the SmCa\(_{\beta}\) clone was used as template for amplification of the truncated insert by PCR. The forward primer (5'-GGGGATCCATGAAAAATGCGTGCAGGAAACGG-3') was designed to bind to the SmCa\(_{\beta}\) clone, starting from nucleotide 142. This forward primer contains a start codon, a Kozak
sequence and a BamHI restriction site. The reverse primer was the same one used to amplify the full-length sequence. These PCR products were inserted into pXOOM and transformed into *E. coli*. Clones were sequenced to verify the deletion and to detect possible PCR errors.

**Electrophysiology**

Whole-cell recordings were obtained at room temperature 24 hours following transfection using an Axopatch 200B (Molecular Devices). Cell capacitance was 12–25 pF. Series resistance was compensated by 70%. Voltage pulses from -20 mV to +70 mV were delivered in 5 mV increments from a holding potential of -80 mV. Data were acquired at sampling intervals of 50 μs and filtered at 5 kHz during acquisition. The pipette solution contained (mM): cesium methane sulfonate (110), HEPES (10), EGTA (9), Mg2+-ATP (5); pH (CsOH) 7.3, with variations, as noted. The bath solution contained (mM): CaCl2 (10),

**Figure 5**

Intracellular free Na⁺ does not block Caᵥ2.3 channels expressed alone or co-expressed with SmCaᵥβ or Caᵥβ2a but induces SmCaᵥβ-dependent run-down of Caᵥ2.3 currents. Current-voltage relationships from HEK cells expressing Caᵥ2.3 subunits alone (A), co-expressing Caᵥ2.3 and SmCaᵥβ subunits (B) or co-expressing Caᵥ2.3 and Caᵥβ2a subunits (C) in minimal intracellular solution (filled circles) and in intracellular solution containing 5 mM NaCl (open circles). Peak currents generated by voltage steps from -20 mV to +75 mV in 5 mV steps from a holding potential of -80 mV are plotted. Data points represent mean ± s.e.m. (D) Peak current amplitude in the presence of 5 mM internal NaCl relative to peak current amplitude in the absence of NaCl for Caᵥ2.3 subunits alone, co-expressed with SmCaᵥβ subunits or co-expressed with Caᵥβ2a subunits. (E) Relative current amplitude of currents produced by Caᵥ2.3 (filled circles) channels alone and in combination with SmCaᵥβ (open circles) plotted as a function of time in the presence of 5 mM NaCl in the patch-pipette solution. Values are means ± s.e.m, N = 3 – 9.
Figure 6
The N-terminal domain of SmCa\textsubscript{β} is the molecular substrate for Na\textsuperscript{+} and Mg\textsuperscript{2+}-ATP mediated block of Ca\textsubscript{2.3} currents. A. Diagram showing the portion of the N-terminal domain, including a domain rich in aspartic acid and glutamic acid (DE-rich domain), that is removed by the truncation. Acidic residues are in red. B. Current-voltage relationships from HEK cells expressing Ca\textsubscript{2.3} alone (filled circles) and in combination with SmCa\textsubscript{β} (open circles) or N-terminally truncated SmCa\textsubscript{β} (filled triangles). Peak currents generated by voltage steps from -20 mV to +75 mV in 5 mV steps from a holding potential of -80 mV were measured. Data points represent mean ± s.e.m. N = 6–7. C. Voltage-dependence of inactivation (steady-state inactivation) for Ca\textsubscript{2.3} alone co-expressed with SmCa\textsubscript{β} (filled circles) or with its N-terminally truncated version (open circles). Lines represent fits to the Boltzmann function. D. Relative peak current amplitude of currents produced by Ca\textsubscript{2.3} channels in the presence of internal 5 mM Mg\textsuperscript{2+}-ATP co-expressed with the wild type SmCa\textsubscript{β} subunit (filled circles) or with the N-terminally truncated SmCa\textsubscript{β} subunit (open circles) plotted as a function of time. Currents were normalized with respect to those measured immediately after rupturing the patch membrane (t = 0). Data points represent mean ± s.e.m. N = 5. E. Relative current amplitude of currents produced by Ca\textsubscript{2.3} channels in the presence of internal 5 mM NaCl co-expressed with the wild type SmCa\textsubscript{β} subunit (filled circles) or with the N-terminally truncated SmCa\textsubscript{β} subunit (open circles) plotted as a function of time. Currents were normalized with respect to those measured immediately after rupturing the patch membrane (t = 0). Data points represent mean ± s.e.m. N = 10–13. F. Comparison between the relative current amplitudes produced by Ca\textsubscript{2.3} channels co-expressed with SmCa\textsubscript{β} (in black) or with its N-terminally truncated version (in grey), in the presence of intracellular 5 mM Mg\textsuperscript{2+}-ATP (left), or in the presence of 5 mM NaCl (right), after 2.5 minutes of establishing the whole-cell configuration. Asterisks denote statistical difference with respect to current amplitude for Ca\textsubscript{2.3} co-expressed with wild type SmCa\textsubscript{β} (Student t test, p < 0.05).
TEA-Cl (160), HEPES (10), EGTA (0.1): pH (TEA-OH) 7.4. Patch pipettes were pulled from borosilicate glass and fire polished before each experiment. To ensure a fast dialysis of the intracellular compartment, only pipettes with resistances between 0.8 and 1.2 MΩ were used. Membrane seals were obtained by applying negative pressure. All experiments were performed at room temperature (22°C). The voltage-dependence of steady state inactivation was determined by measuring the peak current evoked with a depolarising pulse to elicit the maximum current as a function of the voltage of a preceding 1.5 s pre-pulse test (between -110 and +20 mV). Steady-state inactivation curves were fitted by a sigmoid (Boltzmann) distribution of the form:

\[ F(V) = \frac{I_{\text{max}}}{1 + \exp \left( \frac{V_{0.5} - V}{K} \right)} \]

Where \( I_{\text{max}} \) is the maximal current, \( V \) is the pre-pulse voltage, \( K \) is the slope factor and \( V_{0.5} \) is the voltage at which inactivation is half-maximal.

To study inactivation as a function of time, the decaying phases of the inward currents evoked by a test pulse to +20 or +30 mV were fitted to a double exponential equation of the form: \( I(t) = I_0 + I_1 \exp(-t/\tau_1) + I_2 \exp(-t/\tau_2) \), where \( \tau_1 \) and \( \tau_2 \) represent the fast and the slow time constants of inactivation, respectively, \( I_1 \) and \( I_2 \) represent the relative contribution of each component to inactivation and \( I_0 \) is the offset. To assess the rate of recovery from inactivation from the closed state, a two-pulse protocol was used. Two test pulses to +20 or +30 mV were separated by a recovery step to -80, -100 or -120 mV for varying amounts of time (from 10 ms to 1 second). The length and voltage of the first test pulse was adjusted accordingly for each particular channel combination. For example, currents produced by Cav2.3 channels alone were maximal at approximately +30 mV and inactivated completely within 100 ms, whereas currents produced by Cavα,β were maximal at approximately +20 mV and required several seconds to inactivate fully. In the latter case, recovery from inactivation was measured using Ba\(^{2+}\) instead of Ca\(^{2+}\) as the charge carrier (Ba\(^{2+}\) currents inactivate faster than Ca\(^{2+}\) currents produced by Cav2.3 [45]), thereby decreasing the need to use an excessively long depolarising first pulse protocol, which could compromise cell viability. The currents evoked by the second pulse of this double-pulse protocol were normalized to the currents produced by the first pulse and plotted against the duration of the interpulse interval.

**Statistical analyses**

Statistical comparisons were carried out using the Student t-test. Data are presented as means ± s.e.m. Number of repeats is indicated in parentheses.

**Authors’ contributions**

VSR carried out the molecular and electrophysiological studies, made substantial contributions to conception and experimental design, and drafted the manuscript. TS provided the HEK cell line stably transfected with Ca\(_{2.3}\) and made a significant contribution to the interpretation of the data. RMG carried out molecular work and contributed to experimental design as well as to the writing of the manuscript. All authors read and approved the final manuscript.

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