TRIC-B channels display labile gating: evidence from the TRIC-A knockout mouse model

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Abstract Sarcoplasmic/endoplasmic reticulum (SR) and nuclear membranes contain two related cation channels named TRIC-A and TRIC-B. In many tissues, both subtypes are co-expressed, making it impossible to distinguish the distinct single-channel properties of each subtype. We therefore incorporated skeletal muscle SR vesicles derived from Tric-a-knockout mice into bilayers in order to characterise the biophysical properties of native TRIC-B without possible misclassification of the channels as TRIC-A, and without potential distortion of functional properties by detergent purification protocols. The native TRIC-B channels were ideally selective for cations. In symmetrical 210 mM K⁺, the maximum (full) open channel level (199 pS) was equivalent to that observed when wild-type SR vesicles were incorporated into bilayers. Analysis of TRIC-B gating revealed complex and variable behaviour. Four main sub-conductance levels were observed at approximately 80% (161 pS), 60% (123 pS), 46% (93 pS), and 30% (60 pS) of the full open state. Seventy-five percent of the channels were voltage sensitive with Po being markedly reduced at negative holding potentials. The frequent, rapid transitions between TRIC-B sub-conductance states prevented development of reliable gating models using conventional single-channel analysis. Instead, we used mean-variance plots to highlight key features of TRIC-B gating in a more accurate and visually useful manner. Our study provides the first biophysical characterisation of native TRIC-B channels and indicates that this channel would be suited to provide counter current in response to Ca²⁺ release from the SR.

Further experiments are required to distinguish the distinct functional properties of TRIC-A and TRIC-B and understand their individual but complementary physiological roles.

Keywords Sarcoplasmic reticulum · Ion channels · Excitation–contraction coupling · Ca²⁺ release

Introduction

The TRIC channel is an intracellular ion channel that forms a homo-trimeric assembly with a bullet-shaped structure [20, 32]. Two TRIC subtypes, namely TRIC-A and TRIC-B, are found on endo/sarcoplasmic reticulum (ER/SR) and nuclear membranes in animal cells. TRIC-A is especially
abundant in excitable tissues, such as cardiac and skeletal muscle, while TRIC-B is found in most tissues [32]. We have previously demonstrated that recombinant, purified TRIC-A and TRIC-B proteins behave as monovalent cation channels when reconstituted into artificial membranes and that they display many similar biophysical properties to the native SR K+ channels that are observed when SR vesicles from skeletal and cardiac muscle are incorporated into bilayers [7, 15, 16, 26, 27].

The physiological role of TRIC channels is not yet fully established. There is good evidence, however, to support the hypothesis that they act as a counter current pathway, enabling sufficient monovalent cation flux to buffer the loss of positive charge that results during SR Ca2+-release events [30, 32]. Much of this evidence comes from studying the pathology of the Tric-knockout mouse models [30–33]. In the Tric-b-knockout mouse, the alveolar epithelial cells exhibit inadequate Ca2+-release from overloaded ER stores, and this causes the mice to die in respiratory failure following birth [30]. In mutant mice lacking both TRIC-A and TRIC-B, the cardiac SR becomes swollen and cardiac arrest is observed after embryonic day E9.5 [32]. Although the Tric-a-knockout (Tric-a-KO) mouse is viable, the ability of the skeletal muscle cells to respond to stress is defective, and this is revealed by instability in the Ca2+-release process [33]. Moreover, this knockout mouse develops hypertension due to impaired hyperpolarization signalling evoked by Ca2+ spark/release in vascular smooth muscle [31].

The survival of the Tric-a-KO mice provides us with the opportunity of isolating SR membranes from tissues which express only the TRIC-B subtype. This is important because although we have previously investigated the single-channel properties of recombinant TRIC-A and TRIC-B channels purified from cDNA-transfected yeast cells [20], it is well known that purification procedures can alter the functional properties of ion channels [11, 13, 14, 29]. At the same time, it is impossible to properly explore the individual single-channel characteristics of native TRIC-A and TRIC-B channels since they are expressed together in most tissues. Our work with purified TRIC channel subtypes shows that they share certain biophysical properties such as selectivity for monovalent cations, activation by positive holding potentials and complicated sub-conductance state gating behaviour. These similarities make it difficult to definitively distinguish between TRIC-A and TRIC-B channels after incorporating native SR into artificial membranes. In this study, we have therefore isolated SR vesicles from the skeletal muscle of TRIC-A deficient mice. This has allowed us to characterise the basic single-channel properties of TRIC-B channels without interference from the possible co-incorporation of TRIC-A channels into the bilayers. This study provides the first description of the single-channel behaviour of native mouse TRIC-B channels. In future studies, it will be important to investigate the functional differences between TRIC-B and TRIC-A channels as such experiments will provide essential data for understanding their complementary yet distinct physiological roles.

Materials and methods

Preparation of sucrose gradient fractions

As previously described, a discontinuous sucrose gradient was used to separate sarcolemmal membrane vesicles from the various SR membrane fractions, including light SR (LSR) and heavy SR (HSR) from wild-type (WT) mouse skeletal and cardiac muscle and from Tric-a-KO mouse skeletal tissue [23, 24]. For each tissue, the various membrane fractions were aliquoted and stored at −80 °C.

Planar phospholipid bilayer techniques

Recombinant TRIC-B channels or LSR or HSR vesicles from WT and Tric-a-KO tissues were reconstituted into planar phosphatidylethanolamine lipid bilayers as described previously [24]. Recording solutions were symmetrical, unless otherwise indicated, and contained 210 mM KPIPES, 10 μM free Ca2+, pH7.2. Briefly, after forming a gradient of K+ between the two compartments, recombinant proteins or SR vesicles were added to the cis (cytosolic) side of the bilayer. After channel incorporation, the cis chamber was perfused with fresh KPIPES solution to restore symmetrical conditions. The trans (luminal) chamber was held at ground, and the cis chamber was clamped at potentials relative to ground. Experiments were performed at room temperature (22±2 °C). The free [Ca2+] and pH of the solutions were maintained constant during the experiment and were determined using a Ca2+ electrode (Orion 93-20) and a Ross-type pH electrode (Orion 81-55) as previously described [24]. After addition of 10 mM EGTA to the cis chamber without further addition of Ca2+, the approximate free [Ca2+] was calculated using the program MaxChelator (www.stanford.edu/~cpatton/maxc.html) to be <1 nM.

Criteria for defining TRIC-B channels

Addition to the cis chamber, of LSR or HSR vesicles from Tric-a-KO tissue, in the presence of a KCl gradient, led to the rapid incorporation of both cationic and anionic channels. In symmetrical solutions of KPIPES, only two distinct types of K+ current fluctuations were routinely observed: (1) those we describe as TRIC-B currents (observed in 100 % of bilayer experiments), characterised by full open channel conductance of 200 pS but always observed to visit multiple sub-conductance states as described below; (2) those
corresponding to currents through RyR channels (single-channel conductance of approximately 700 pS). RyR channels incorporated less frequently into the bilayer when LSR rather than HSR fractions were used, and for this reason, only TRIC-B channels from the LSR fraction was used. Supplementary Fig. 1 illustrates a typical recording where both an RyR and a TRIC-B channel from the LSR have incorporated into the bilayer demonstrating that the single-channel characteristics of these two channels are very different and could not be confused. We only used those recordings where no RyR channels were observed and where no leak in the baseline was detected. It is possible that there may be contamination of our SR membrane fractions by sarcolemmal membrane vesicles. However, when skeletal muscle sarcolemmal vesicles are fused into bilayers, fusion events are slower and less reproducible in terms of the type of current fluctuations that are observed. Fusion of sarcolemmal membranes with a bilayer occasionally led to the appearance of a Ca\(^{2+}\)-activated K\(^+\) channel of conductance of approximately 250 pS in symmetrical 210 K\(^+\) solutions, but we did not observe this channel after fusion of LSR membrane vesicles with the bilayer. If this channel had incorporated during fusion of LSR vesicles, we would have been able to distinguish it from a TRIC channel on the basis of conductance, sub-conductance gating behaviour and Ca\(^{2+}\) dependence. The Ca\(^{2+}\) dependence of the Ca\(^{2+}\)-activated K\(^+\) channel is well known (for example [12, 17]). In our hands, the Ca\(^{2+}\)-activated K\(^+\) channel that we observe from the sarcolemmal fraction is closed by reducing the cis free [Ca\(^{2+}\)] to 0.1 μM (results not shown), whereas TRIC-B is not regulated by Ca\(^{2+}\) in this fashion (see below).

Data acquisition and analysis of single-channel recordings

Single-channel recordings were digitised at 100 kHz and recorded on a computer hard drive using WinEDR 3.05 software (John Dempster, Strathclyde University, UK). Current–voltage relationships were constructed using current amplitudes, which were measured using manually controlled cursors in WinEDR 3.05. The conductance values were obtained by linear regression (Graphpad Prism 4, La Jolla, USA). We consistently observed four sub-conductance states at +30 and −30 mV. At higher holding potentials or improved resolution of the single-channel events, it is likely that more sub-conductance states would be detected. Using our initial sub-conductance state amplitude measurements, single-channel current fluctuations were idealised using the segmental k-means (SKM) algorithm [21] in the QuB software suite (State University of New York, Buffalo, USA). Before idealisation, traces were digitally filtered (Gaussian Filter with cut-off frequency of 1 kHz) and re-sampled at 10 kHz. A dead time of 600 μs was imposed on each idealisation. A six-state Markov model was constructed with identical fully connected topology, where states consisted of the closed, open and four sub-conductance states using our initial estimates of the current amplitudes of the conductance states. After building the model, the Viterbi algorithm [2, 5] was applied in order to estimate the best fit to our data based on the log-likelihood function value. The parameters of the model were then used to plot amplitude probability density functions which were superimposed on the all-points amplitude histogram. The idealisations obtained with the SKM method were also superimposed on the single-channel traces to visually examine the accuracy of the method. The probability of dwelling in each of the states (closed, open and four sub-conductance states) was determined from the SKM idealisation of continuous recordings of 2–3 min at holding potentials of +30 and −30 mV. To check whether a single channel only was present in the bilayer, high positive holding potentials were applied for short periods of time to maximally activate the channel at the end of the experiment.

Average current, I, was calculated using the following equation:

\[
I = P_{\text{full}}I_{\text{full}} + P_{S1}I_{S1} + P_{S2}I_{S2} + P_{S3}I_{S3} + P_{S4}I_{S4}
\]

where \(P_{\text{full}}, P_{S1}, P_{S2}, P_{S3}, \) and \(P_{S4}\) are the mean probabilities of the channel dwelling in each conducting state (full open and S1–S4) and \(I_{\text{full}}, I_{S1}, I_{S2}, I_{S3}, \) and \(I_{S4}\) are the mean current amplitudes for the corresponding conductance levels. Average currents were computed for recordings containing a single-channel gating at holding potentials of +30 and −30 mV.

Mean-variance histograms were computed using the method described by Patlak [18] and implemented in C and Python, from traces digitally filtered (Gaussian Filter) at 800 Hz and using at least 1 min of continuous recording. Briefly, a window of given length was stepped along the data, producing a pair of mean and variance values for each window location. These co-ordinate pairs were then binned in two-dimensions producing the histogram. For the histograms presented here, a window length of 10 ms was used.

SDS-polyacrylamide gel electrophoresis and immunoblot analysis

To examine the distribution of TRIC subtypes in the mouse muscle SR, total microsomes were prepared from WT mice and then separated by discontinuous sucrose density gradient centrifugation to yield the surface membrane (sarcolemma and transverse tubule-rich), longitudinal SR-rich (LSR) and junctional SR-rich (HSR) fractions as previously described [23]. Various fractions were subjected to SDS-polyacrylamide gel electrophoresis and Western blotting using previously described methods [20, 32] as required. Antibodies to TRIC-A, TRIC-B, and Junctophilin type 1 (JP1) were produced.
inhouse and the antibody for RyR was purchased from Thermo Fisher Scientific Inc. (Rockford, USA).

Statistics

Data are expressed as mean±SD, where \( n=3 \), or mean±SEM, where \( n \geq 4 \). Differences between mean values were assessed using a Student’s \( t \) test. A \( p<0.05 \) was taken as significant.

Materials

All chemicals were purchased from VWR (Poole, UK) or Sigma-Aldrich (Dorset, UK). All solutions were prepared in MilliQ deionised water (Millipore, Harrow, UK), and those for use in bilayer experiments were filtered through a membrane with 0.45-μm pore diameter (Millipore, Harrow, UK).

Results

Figure 1 shows representative recordings of the \( K^+ \) current fluctuations that are observed when LSR vesicles isolated from the heart and skeletal muscle of WT mice are incorporated into artificial bilayers in symmetrical solutions of 210 mM KPIPES. These single-channel events look similar to those obtained from cardiac or skeletal SR vesicles of other species, including rabbit, sheep, dog, and bullfrog [3, 4, 7–10, 15, 16, 20, 26, 28]. Channel gating was characterised by openings to relatively long-lived full open states and by transitions to other sub-conducting open states that often look ‘messy’ or noisy. The full open state was 211±17 pS (SD; \( n=6 \)) for channels derived from skeletal SR and 200±5 pS (SD; \( n=4 \)) for those from cardiac SR. Since we know that both TRIC-A and TRIC-B are expressed in striated SR, we cannot definitively assign any channel opening as belonging to TRIC-A or TRIC-B. We have therefore used two different approaches that guaranteed the absence of the TRIC-A isoform. We have incorporated (1) the skeletal muscle LSR membrane fraction from Tric-a-KO mice or (2) recombinantly expressed, detergent-purified mouse TRIC-B into bilayers in symmetrical 210 mM KPIPES.

In rabbit skeletal muscle, TRIC-A is immunochemically detected in the nuclear and SR membranes [32]. Figure 2a shows the distribution of TRIC subtypes in the mouse muscle membrane fractions. As expected from previous observations, both TRIC subtypes were slightly enriched in the HSR, but were distributed throughout the SR fractions (Fig. 2b). At the same time, because of the similar protein compositions of the microsomal preparations from WT and Tric-a-KO mice (Fig. 2c), SR protein expression seems roughly normal in Tric-a-KO muscle. Indeed, Western blot analysis detected regular expression of TRIC-B, RyR and junctophilin type 1 in Tric-a-KO muscle (Fig. 2d). The SR preparations from Tric-a-KO mice therefore appear to provide an ideal model system from which to examine native TRIC-B channels.

When Tric-a-KO skeletal LSR vesicles were reconstituted into bilayers in order to observe native TRIC-B channel gating in isolation, we observed channel gating that was always characterised by openings to several sub-conductance states. In the event of a simultaneous incorporation of RyR channels, the record was discarded. A representative example of a native TRIC-B channel recording is illustrated in Fig. 3a. The full open channel conductance of native TRIC-B channels was approximately 200 pS and was consistently

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Fig. 1 Native TRIC channel currents from mouse cardiac and skeletal LSR. Examples of \( K^+ \) current fluctuations recorded after the incorporation of WT mouse cardiac (a) and skeletal (b) LSR membrane vesicles in symmetrical solutions of 210 mM KPIPES. Channel openings are upward deflections, and \( O \) and the black arrow indicate the open and closed channel levels, respectively.
observed in 34/34 experiments. This was similar to the full open state of the channels derived from WT SR (Fig. 1). Channel gating was also characterised by rapid transitions between multiple sub-conductance states. Many of these events were too brief to resolve, and so we have only measured those levels that could be detected manually. We found that the channels dwelt in four predominant sub-conductance levels that were approximately 80, 60, 46, and 30% of the full open state. These are detailed in Table 1. We predict that further sub-conductance states will be observed as the resolution of the single-channel events is improved. The corresponding current–voltage relationships of the full and the four sub-conductance levels detected are shown in Fig 3b for the channel shown in Fig. 3a.

<table>
<thead>
<tr>
<th>% of full open state</th>
<th>Full open</th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
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<tr>
<td>Mean±SEM (pS)</td>
<td>199±2</td>
<td>161±2</td>
<td>123±1</td>
<td>93±2</td>
<td>60±2</td>
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<tr>
<td>n</td>
<td>34</td>
<td>18</td>
<td>24</td>
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Table 1 Native TRIC-B conductance values for the full and the four main sub-conductance levels
When we incorporated recombinantly expressed, detergent-purified TRIC-B channels into bilayers, similar current fluctuations with multiple sub-conductance state openings to those observed in Figs 1 and 3 were observed with the exception that the channels did not appear to open to the full open state. The data are shown in Supplementary Fig. 2. We concluded that the expression system and/or purification procedure may have led to changes in protein folding and altered functional behaviour of TRIC-B channels so that they rarely open to the full open state. We therefore focused exclusively on using the native TRIC-B channels obtained from Tric-a-KO tissue in order to characterise the key aspects of TRIC-B gating.

To investigate whether TRIC-B is selective for cations, experiments were performed in a KCl gradient (810 mM K+, 600 mM Cl⁻ cis; 210 mM K+ trans, Fig 4a). A parallel shift to the left but no change in single-channel conductance was observed. The maximum conductance of the fully open channel in KCl gradient was 212±8 pS (SD, n=6), and the reversal potential ($E_{rev}$) was $−37.7$ mV. These values are similar to the calculated value for a channel ideally selective for cations, where $E_{rev}$ is $−34$ mV.

The free cytosolic $[\text{Ca}^{2+}]$ was maintained at 10 μM to be consistent with our previous experiments with purified and native TRIC and RyR channels. We have previously reported that purified TRIC-B channels were activated by raising cytosolic $\text{Ca}^{2+}$ from 10 to 100 μM [20]. To examine whether cytosolic $\text{Ca}^{2+}$ is a requirement for TRIC-B activation, we lowered the free $[\text{Ca}^{2+}]$ to below 1 nM by addition of a high concentration of EGTA. No significant change in Po was observed. The mean data is shown in Fig. 4b. These experiments demonstrate that, although previous experiments show that TRIC-B channels exhibit sensitivity to changes in cytosolic $\text{Ca}^{2+}$ above 10 μM, $\text{Ca}^{2+}$ is not an absolute requirement for channel activation as the channels still open at free $\text{Ca}^{2+}$ concentrations that are well below the resting levels found in cells.

Detailed analysis of TRIC-B gating was conducted only for recordings where a single channel was gating in the bilayer ($n=8$). Identification of sub-conductance levels was extremely difficult in records containing multiple channels. We found that a key feature of native TRIC-B channel gating was the voltage dependence. A representative example of a TRIC-B single-channel recording at holding potentials of +30 and $−30$ mV is shown in Fig. 5a. TRIC-B Po was significantly reduced at negative holding potentials. The probability of dwelling in the full open channel level was 0.041±0.011 at +30 mV but only 0.009±0.007 at $−30$ mV ($n=8$; SEM; $p<0.05$).

Examples of typical sub-conductance gating of single TRIC-B channels and the corresponding amplitude histograms at +30 and $−30$ mV are illustrated in Fig. 6. For each trace, a 200-ms section of the recording is shown on an expanded time scale so that transitions to the four main sub-conductance states are easily visualised. It should be noted that TRIC-B openings at negative potentials were sporadic and that channel gating consisted primarily of transitions to sub-conductance states that did not reach the full open channel level as shown in Fig. 6e.

Mean data of the probabilities of dwelling in the full, the closed and the four sub-conductance states (S1, S2, S3 and S4) at holding potentials of +30 and $−30$ mV for the eight single channels are compared in Fig. 7a. Grouping the data in this manner suggests that the channels spend longer in the sub-conductance open states (especially S4) than in the full open state and that there was virtually no voltage dependence of gating. However, this was not what we observed for most channels. Further inspection of the data showed that two channels were heavily weighting the data and so we excluded them from the group analysis. Mean data was then obtained for the six remaining channels (channels 1–6;
Fig. 7b), and the values for the two outliers, channels 7 and 8, were plotted individually.

This simple analysis clearly demonstrates that the gating of TRIC-B is variable and complex and that there are at least three distinct patterns of gating. The first and most regularly observed pattern \((n=6/8)\) was seen in channels which, at positive potentials, exhibit a higher probability of dwelling in the full open state than in the subconductance states. These channels also exhibited strong voltage-dependent gating and were more open at +30 than −30 mV. Importantly, the main determinant of the voltage dependence of open probability \((\text{Po})\) was the reduction in the time spent in the full open state. The mean data for this group of channels is shown in Fig. 7b.

A second pattern of gating was observed in one of the outlying channels (channel 7). This channel was voltage dependent but dwelt for longer in the sub-conductance states (especially S4) than in the fully open state at both +30 and −30 mV (Fig. 7c). In this channel, the probability of dwelling in all of open states (full and sub) was reduced at −30 mV in comparison to +30 mV. A third type of gating was observed in channel 8. Like channel 7, channel 8 also dwelt for longer in the sub-conductance states (especially S4 and S3) than in the fully open state, but it was voltage independent and had very high overall Po (Fig. 7d). Sections of the single-channel record of this particular channel and the corresponding amplitude histograms illustrating the predominant gating in the lower sub-conductance states are shown in Fig. 8. It can be seen that the amplitude histogram distributions are very different to those shown in Fig. 6.

The rapid transitions between multiple sub-conductance states, together with the fact that TRIC-B channel gating does not conform to a single pattern of gating, makes it difficult to provide a simple and accurate kinetic model that encapsulates the key aspects of TRIC-B gating and that might allow us to quickly distinguish it from TRIC-A. We have therefore plotted mean-variance histograms in order to provide an accurate but also a more easily accessible and visual summary of the important aspects of TRIC-B gating. These are shown in Fig. 9a and b for a typical low Po channel belonging to the set of 6 channels grouped together in Fig. 7b. In Figure 9c and d, the mean-variance plots for outlier channel 8, which exhibited high Po, voltage
independence and predominance of gating in sub-states S3 and S4 are shown for comparison.

The colour density of the plots is proportional to the number of samples binned for a given mean-variance pair, and, therefore, if colour density is high (yellow) for a particular current amplitude, this reflects a high number of transitions to and from open/closed states of that current amplitude. Regions with a constant (generally low) variance and a high sample density may correspond to discrete conductances of the TRIC-B channel such as those identified and described in Table 1. Arcs through high-variance regions often correspond to transitions between states and high density in these regions indicates a large number of transitions.

For the low Po, voltage-dependent channel at +30 mV (Fig. 9a), the points are clustered closely around the closed state, with the region of next highest density (and low variance) being the full open state. Arcs through regions of higher variance correspond to channel transitions to and from the various sub-conductance states and reflect the comparatively brief dwell times in those states. At negative potentials (Fig. 9b), the majority of time is spent in the closed state, and it is clear that there are no detected transitions to the fully open state of TRIC-B. On the bottom, an expanded section of the trace is illustrated (100 ms). In this case, the channel appears to gate only to the sub-conductance states S3 (blue) and S4 (red).
Transitions to and from the small conductance sub-states (S4 and S3) give rise to areas of particularly high variance for a given mean current. The large areas of high variance between the black and red and blue arrows indicate high numbers of short-lived events and rapid transitions between the closed states and S3 and S4. Likewise, at $-30 \text{ mV}$ (Fig. 9d), the channel is still opening to a high Po level and a considerably large proportion of the time that the channel is open is spent switching between the full open and sub-conductance states rather than dwelling at any one particular state for long periods of time.

### Discussion

By knocking out expression of TRIC-A, we have been able to study the single-channel behaviour of native mouse skeletal TRIC-B channels. This has allowed a detailed study of the biophysical characteristics of this channel without misclassifying any single-channel current fluctuations as TRIC-A events. Unexpectedly, we find that the full open state conductance of native mouse TRIC-B ($199 \text{ pS}$) is significantly higher than our measurements of the highest conductance open events of purified recombinant TRIC-B (which varies between $140–160 \text{ pS}$ under identical recording conditions (see Supplementary Fig. 2)). Instead, the full open state conductance of native TRIC-B is similar to our reported values for purified recombinant TRIC-A ($192 \text{ pS}$) \[20\]. It is also similar to the highest conductance open state that we observe when we incorporate WT mouse SR into bilayers ($200 \text{ pS}$).

We attribute the apparent lower conductance of purified TRIC-B to modification of the gating behaviour of the channel by the purification procedure and/or the use of a non-mammalian expression system. It appears that the purified TRIC-B channels gate only in sub-conducting open states. We used CHAPS as the detergent in this study but have previously developed protocols using two other detergents ($n$-dodecyl beta-D-maltopyranoside \[20\] and...
n-decyl-beta-D-maltopyranoside (unpublished observations)), but in all cases, recovery of functional ion channels was problematic and we observed only low conductance events ($\leq 160$ pS). Aside from the problems associated with protein purification, *E. coli* and yeast cells may not be appropriate expression systems for TRIC channels and may lead to incorrect protein folding, which does not support normal channel gating behaviour. These experiments highlight the need to study native channels.

Our experiments also revealed that TRIC-B gates to a number of sub-conductance gating states. With the resolution of single-channel events possible at the holding potentials used (+30 and −30 mV), we identified four frequently visited sub-conductance gating states. With improved resolution, it is likely that additional sub-conductance levels will be revealed. We were not able to hold at higher potentials for long periods of time in order to improve the signal/noise ratio as this frequently broke the bilayers. It was usual for multiple channels to incorporate into the bilayers, but in eight experiments, we found that only single channels were active in the bilayer and we used these experiments to analyse the gating behaviour of TRIC-B channels in more detail. Previous reports suggest that SR K$^+$ channels exhibit a single sub-conductance state of approximately 60–70 % of the fully open state. This sub-state has often been described as messy or noisy and there are quite wide variations in the reported amplitudes of the sub-conductance state [6, 10, 15, 19, 22, 26, 27]. We suggest that the noisy gating of this reported 60–70 % sub-conductance state actually reflects the rapid transitions between multiple sub-conductance states of the TRIC channels (see Figs. 6 and 8 for examples). When normal or WT SR is incorporated into bilayers, we assume that both TRIC subtypes could be incorporated. It is feasible that most previous recordings were of TRIC-A channels, since estimated TRIC-A levels are roughly 10 times more than TRIC-B levels in cardiac and skeletal SR preparations based on quantitative reverse transcriptase polymerase chain reaction (RT-PCR) and Western blotting data (D. Y. and H. T., unpublished observations). However, future work investigating the biophysical properties of native TRIC-A channels is required to address this issue.

Our experiments suggest that we may not be able to distinguish between TRIC-A and TRIC-B on the basis of single-channel conductance measurements, and this led us to attempt to define the hallmark features of TRIC-B gating. However, although there are major trends in gating

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**Fig. 8** Typical current fluctuations through the voltage-independent channel. a, b Representative sections of the voltage-independent, high Po TRIC-B channel (channel 8) recording at +30 and −30 mV. The black arrow indicates the closed channel level (dashed black line) while the full open conductance is indicated by O (pink). Sub-conductance states, in order of increasing current amplitude, are represented by S4 (red), S3 (blue), S2 (green) and S1 (yellow). Expanded sections (200 ms) show the sub-conductance gating behaviour with SKM idealization overlaid (orange). c, d All-points amplitude histograms of the portions of trace shown in a and b, respectively. Gaussian distributions computed in SKM fit for the closed, the open and Sn states, are superimposed and coloured as above. The thick black trace represents the sum of Gaussians and the combined fit of the model’s state amplitudes to the filtered data.
behave it is evident that TRIC-B gating is extremely variable and that a single unified model of gating would not be adequate. In terms of hallmark features, voltage dependence was the most obvious trait, and yet this was not observed for all channels. The majority of channels (7/8) were strongly voltage dependent with very low Po values at $-30 \text{ mV}$, but one channel was clearly not regulated by voltage and exhibited high Po at both $\pm 30 \text{ mV}$. Another prevalent feature was the dominance of the full open state at $+30 \text{ mV}$; the Po of six in eight channels was primarily determined by openings to the fully open state, and only two of the eight channels opened predominantly to the sub-conductance states. It is possible that the channels that appear to be outliers (shown in Fig 7c and d) may simply represent examples of the wide range of gating behaviour that occurs physiologically.

**Fig. 9** Representative mean-variance histograms showing the sub-conductance gating behaviour exhibited by TRIC-B channels. The colour density is proportional to the number of samples binned for a given mean-variance pair. The regions with a constant (generally low) variance and a high sample density may correspond to discrete conductances of the TRIC-B channel. Arcs through high-variance regions often correspond to transitions between states and high density in these regions indicates a large number of transitions. The arrows indicate the closed (black), open (pink) and sub-conductance S1–S4 (yellow, green, blue and red, respectively) state levels that were measured in the corresponding single-channel traces. **a, b** Typical histograms for a channel belonging to the set of channels grouped together in Fig. 7b (channels 1–6). These channels were characterised by voltage-dependent gating and higher Po in the full open state than in the sub-conductance states. At $+30 \text{ mV}$, the channel is closed (0 pA) most of the time with some brief transitions to various sub-states. These transitions manifest themselves as arcs through high-variance regions; however, because they are relatively infrequent, the individual sub-conductance states are not defined by clear high-density regions. For this channel, it can be seen that the majority of the openings are to the full open state. Conversely, at $-30 \text{ mV}$, the samples are almost exclusively distributed around the closed state, with a small bias towards S4 (red arrow), the lowest conductance open state. There are no openings to the full open state. **c, d** Voltage-independent gating of channel 8. The channel dwells for a noticeably longer proportion of time in sub-conductance open states than the full open state at both $+30$ and $-30 \text{ mV}$. The samples in the mean-variance plot are clustered with similar density around regions with low variance representing mean currents that correspond to sub-conductance states. Rapid gating between closed, S4 and S3 states gives rise to a slightly higher variance region for a given mean than would be expected for long sojourns in a specific conductance state. The density of samples around the open states is much higher than in **a and b**, corresponding to a much higher overall Po.
We have used mean-variance plots as a method of highlighting the important aspects of TRIC-B gating in a snapshot. This approach allows us to visualise not only the approximate Po of the channel but also the preponderance of sub-conductance state gating and the frequency of transitions between certain states. The plots demonstrate how the usual gating of TRIC-B that is observed in 75% of the channels (six of eight) is dominated by openings to the fully open state at +30 mV but by sub-state gating at −30 mV (Fig. 9a, b). Similarly, it is easy to observe the higher Po and frequent transitions between sub-conductance states at both +30 and −30 mV for the channel data shown in Fig. 9c and d. Our goal is to use this new method of visualising TRIC channel gating to compare and contrast the gating behaviour of TRIC-A and TRIC-B channels. We will also be able to assess how regulators of TRIC channels modify gating without building complicated kinetic gating models that are subject to error because of the poor resolution of the brief sub-conductance events.

As yet, we do not have opportunity to work with native TRIC-A channels since the Tric-b-KO mouse is not viable. Figure 10, however, illustrates how mean-variance plots can be an effective method of highlighting differences in single-channel conductance, gating behaviour and the prevalence of variance at these current amplitudes. The current amplitude of the full open state of RyR is much higher than that of TRIC-B, and this leads to higher overall variance in regions that correspond to rapid transitions between states. RyR gating under the experimental conditions of the recording (symmetrical 210 mM K+, 10 μM free Ca2+) is typified by rapid transitions between the closed and fully open channel levels that are too brief to resolve and which are truncated by filtering. This gives rise to histograms with arcs of high density and high variance. Qualitatively, this type of gating behaviour is clearly different to the sub-conductance gating activity of TRIC-B, where longer-lived sub-conductance states can be distinguished from transitions between states by their lower overall variance and discrete regions of higher density. For both TRIC-B and RyR, the holding potential was +40 mV. The data was sampled at 100 kHz and filtered at 800 Hz using 65–80 s of continuous recording.

### Table 2: Average currents for the three groups of channels and the percent reduction of the average currents observed at −30 mV

<table>
<thead>
<tr>
<th>Group</th>
<th>Average current (pA)</th>
<th>% reduction at −30 mV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>+30 mV</td>
<td>−30 mV</td>
</tr>
<tr>
<td>Channels 1–6</td>
<td>0.320</td>
<td>0.051</td>
</tr>
<tr>
<td>Channel 7</td>
<td>0.963</td>
<td>0.196</td>
</tr>
<tr>
<td>Channel 8</td>
<td>1.598</td>
<td>1.939</td>
</tr>
</tbody>
</table>

### Table 3: Contribution of sub-conductance state gating to the average current for the three groups of channels

<table>
<thead>
<tr>
<th>Group</th>
<th>Average current attributed only to sub-state gating (pA)</th>
<th>% of total current</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>+30 mV</td>
<td>−30 mV</td>
</tr>
<tr>
<td>Channels 1–6</td>
<td>0.075</td>
<td>0.045</td>
</tr>
<tr>
<td>Channel 7</td>
<td>0.782</td>
<td>0.177</td>
</tr>
<tr>
<td>Channel 8</td>
<td>1.310</td>
<td>1.572</td>
</tr>
</tbody>
</table>
of sub-conductance gating states. The figure compares mean-variance plots from single-channel recordings of TRIC-B (top) and RyR (bottom). Note, for example, the high conductance and the very brief openings of RyR channels that are truncated by filtering in comparison to the obvious sub-conductance gating of TRIC-B.

In a cellular context, how would sub-conductance gating contribute to the overall currents that would flow across the SR membrane through TRIC-B? Since we have no reason to suggest that channels 7 and 8 (although atypical for our group of 8 channels) do not form part of the physiological spectrum of TRIC-B channels, how important is their contribution? To answer these questions it is useful to think about average channel current rather than Po in states.

The average TRIC-B current for a channel can be calculated from the current amplitude of the full and sub-conductance states and the Po in those states. Table 2 compares the average current measurements for the different groups of channels (channels 1–6, channel 6 and channel 7). These data show that channels 1–6 have low current overall in comparison to channels 7 and 8, and exhibit an approximately 80 % reduction in current at negative voltages. Channels 7 and 8 would provide most of the current across the membrane at positive potentials if this group of channels was representative of the TRIC-B channel population in the SR or ER of a cell. Channel 8 would also provide significant current at negative holding potentials and markedly alter the overall voltage dependence of the eight channels as a group.

Table 3 shows the contribution of the sub-conductance states to the average channel current. Sub-conductance state gating was low for channels 1–6, contributing only 23 % of the overall current at +30 mV, whereas sub-state gating accounted for approximately 80 % of the average current for channels 7 and 8. For all channels, sub-conductance state gating contributed to approximately 80 % of the total gating at −30 mV. Thinking about the gating of the channels in this manner shows that the outlying channels would contribute significantly to monovalent flux across the membrane and to the overall voltage-dependence of the TRIC-B population of channels. On average, TRIC-B channels would be activated by the development of negative charge in the SR lumen with respect to the cytosol and therefore would be ideally suited to pass counter current to help balance the negative charge that would otherwise develop in the SR during a Ca<sup>2+</sup>-release event. On the other hand, a few voltage-independent channels of high Po such as channel 8 would ensure that there is always a conduit for equilibration of charge across the SR irrespective of voltage. We know very little of how TRIC channels are regulated. It is possible that the variability in TRIC-B gating behaviour that we observe could be due to heterogeneity of channels in regard to post-translational events such as phosphorylation or nitrosylation or maybe due to the presence or absence of a closely associated protein. As such, these may be physiological/pathophysiological mechanisms for stimulating or inhibiting TRIC-B channel opening or inducing sub-conductance state gating.

In summary, we show that native TRIC-B channels are voltage-dependent K<sup>+</sup> channels that exhibit complex and variable gating behaviour. The biophysical properties of TRIC-B indicate that these channels would be well suited to provide monovalent counter current in response to loss of positive charge from the SR during the Ca<sup>2+</sup>-release process. Further studies are required to characterise the single-channel properties of native TRIC-A to identify any key differences between the TRIC channel subtypes that may be important physiologically.

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References


