Ca\(^{2+}\)-activated K channels in parotid acinar cells
The functional basis for the hyperpolarized activation of BK channels

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Abbreviations: BK, a big conductance, Ca\(^{2+}\)-activated K channel; IK1, an intermediate conductance, Ca\(^{2+}\)-activated K channel

Introduction

There are three general classes of Ca\(^{2+}\)-activated K channels. The large conductance, BK channel (K\(_{\text{Ca}1.1, \text{Slo}}\)) is the prominent member of the first group and is activated by depolarized membrane potentials as well as by Ca\(^{2+}\) ions. These channels are ubiquitously expressed throughout the body from secretory cells to the brain. The small conductance, SK, channels (K\(_{\text{Ca}2.1, \text{KCa}2.2\text{ and KCa}2.3}\)) are activated by Ca\(^{2+}\) but not by membrane potential and are widely expressed throughout the central nervous system. The third class has only a single member: the intermediate conductance IK1 (K\(_{\text{Ca}2.3}\)) channel. These channels are also activated by Ca\(^{2+}\) and not by voltage and are less widespread than the other channels. They are quite prominent in red blood cells and T-lymphocytes and in organs involved in salt and fluid secretion including the colon and salivary glands.

IK1 and maxi-K channels are the only K channels expressed in mouse parotid acinar cells.\(^1\)\(^-\)\(^3\) There is extremely little variation in the IK1 protein sequence in different cell types and across species. The mouse parotid IK1 protein is essentially identical to the human variant. In contrast there are several BK channel variants expressed in different cell types and across species. The mouse parotid BK protein is highly homologous with the human form and differs at five, generally conserved, single sites and in the last eight conceptually translated amino acids.\(^1\) There are much larger differences between the mouse parotid BK isoform and the “standard” mouse version (mSlo)\(^4\) that is the common subject of electrophysiological studies using heterologous gene expression. ParSlo does not have the IYF motif at splice site 3 that is present in mSlo. There are differences at splice site 6 including a retained exon in parSlo that contains a premature stop codon.

While BK channel activation is both Ca\(^{2+}\)- and voltage-dependent, these channels can be activated in low (even nominally zero) Ca\(^{2+}\) but this activation requires extremely depolarized potentials—well beyond the physiologically accessible range.\(^4\)\(^,\)\(^5\) Cell Ca\(^{2+}\) levels increase in response to several stimulatory factors but the average, global Ca\(^{2+}\) levels are usually below a few µM, too little to cause significant BK channel activation except at very depolarized potentials. This problem is overcome in many cell types, particularly neurons and muscle cells, by positioning BK channels very near sources of Ca\(^{2+}\) that can produce very high Ca\(^{2+}\) levels in local microdomains.\(^6\)\(^-\)\(^10\)

As noted above, IK and BK channels are expressed in salivary glands including parotid acinar cells. The physiological role of...
these cells in stimulating fluid secretion involves a muscarinic receptor-mediated increase in intracellular Ca\(^{2+}\) and the subsequent activation of the IK1 and BK channels in concert with Ca\(^{2+}\)-activated Cl channels. The increase in Ca\(^{2+}\) in these cells does not exhibit the “hot spots” associated with the microdomains more typical in excitable cells and the maximal global Ca\(^{2+}\) is usually less than 1 \(\mu\)M.\(^{11-14}\) If the BK channels in the parotid cells have the same Ca\(^{2+}\) and voltage sensitivity as their counterparts in other cell types, they would not be activated by these low levels of Ca\(^{2+}\) and yet muscarinic stimulation that results in only modest levels of intracellular Ca\(^{2+}\) clearly activates BK channels.\(^{15}\) Very early studies suggested that a large conductance K channel in salivary glands is activated at low Ca\(^{2+}\) concentrations\(^{16}\) so the BK channels in secretory cells may have properties that differ from those in excitable tissues. We report here our efforts to understand the apparent difference between BK channels in salivary glands and the similar channels in neurons and muscles.

In this study we investigated the Ca\(^{2+}\) dependence of IK1 and BK channel activation in mouse and human parotid acinar cells. Our focus was on intact, whole cell measurements with physiologically relevant solutions. The Ca\(^{2+}\) dependence of parotid acinar IK1 channels was quite similar to the same channels in heterologous expression systems. In contrast we found a considerable difference in the Ca\(^{2+}\) dependence of native parotid BK and previously published studies with heterologously expressed mSlo or hSlo channels. Since, as noted above, the mouse parotid BK channel differs from mSlo and hSlo and the previously published studies utilized different experimental techniques and solutions, we also examined the Ca\(^{2+}\) dependence of heterologously expressed parSlo using the same techniques as for the native cells. Finally, in an effort to understand the functional differences between the expressed channels and those in native parotid cells, we analyzed our results in the context of the Horrigan-Aldrich (HA) model,\(^ {17}\) which has been shown to provide insight into the mechanisms underlying the voltage- and Ca\(^{2+}\)-sensitive properties of BK channels.

**Results**

As described in Introduction, salivary gland acinar cells express two types of Ca\(^{2+}\)-activated K channels: the voltage- and time-independent IK1 channel and a particular, parotid-specific splice variant of BK channels. We have determined the Ca\(^{2+}\) dependence of the activation of these channels in their native context and with physiologically-relevant aqueous solutions.

**Ca\(^{2+}\) activation of IK1 channels.** The results of our investigation of parotid IK1 channels are illustrated in Figure 1. The inset of part A shows the voltage clamp protocol used. Raw currents elicited by this protocol from a parotid acinar cell with 250 nM intracellular Ca\(^{2+}\) are also illustrated in the inset and the currents from the same cell with 20 \(\mu\)M of the IK1 activator DCEBIO added. The current-voltage relations from these data are shown in the main part of Figure 1A and these are zero near the K\(^{+}\) equilibrium potential as expected. Since this concentration of DCEBIO under these conditions produces maximal activation, the ratio between the currents in the presence and absence of DCEBIO allows a reasonable estimate of the fractional IK1
these parotid IK1 channels are quite similar to heterologously expressed IK1 channels (see Discussion).

Ca2+ and BK channel activation.

The Ca2+ dependence of BK channel activation is more complex than that of IK1 channels and examples of this are illustrated in Figure 2. Part A of this figure shows raw BK channel currents recorded at the indicated potentials at -54 mV. These data are fit (see Materials and Methods) by the Hill equation (line) which provides a reasonable description of these data. The apparent $K_f$ and $N$ values obtained from this fit are 360 nM and 3.1, respectively. The $K_f$ values exhibited a small voltage sensitivity (note expanded scale in the inset) ranging from about 340 nM near 0 mV and almost 400 nM at very negative potentials. In contrast the Hill coefficient had little or no voltage dependence (also an expanded scale) with values between about 3.0 and 3.2. The behavior of these parotid IK1 channels are quite similar to heterologously expressed IK1 channels (see Discussion).

Ca2+ and BK channel activation. The Ca2+ dependence of BK channel activation is more complex than that of IK1 channels and examples of this are illustrated in Figure 2. Part A of this figure shows raw BK channel currents recorded at the indicated potentials with the same protocol illustrated in the inset of Figure 1A. Shown are currents from a native parotid acinar cell (upper) and the parotid BK channel variant, parSlo, expressed in CHO cells—note the different range of potentials shown. Both cells in these examples were patched with 80 nM Ca2+. This concentration of Ca2+ was chosen since it is close to the resting concentration in parotid acinar cells.12-14 The relative conductance-voltage relations from these two cells are shown in Figure 2B along with fits

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**Figure 2.** Comparison of the Ca2+ sensitivity of the voltage dependence of native BK channels in parotid acinar cells and the parSlo channel expressed in CHO cells. (A) Raw BK currents recorded from a native cell (upper) and transfected CHO cells (lower) with 80 nM free Ca2+. The voltages used are shown at the right of each current record. (B) Voltage dependence of relative BK channel conductance with 80 nM Ca2+ from a native parotid cell (■) and from a CHO cell transfected with parSlo (○). Lines are fits of the standard Boltzmann equation with apparent valence and $V_h$ values of 1.36 and 46 mV for the native cell data and 2.1 and 164 mV for the heterologously expressed BK channel. (C) Raw BK currents recorded from a native cell (upper) and transfected CHO cells (lower) with 500 nM free Ca2+. The voltages used are shown at the right of each current record. (D) Voltage dependence of relative BK channel conductance with 500 nM Ca2+ from a native parotid cell (■) and from a CHO cell transfected with parSlo (○). Lines are fits of the standard Boltzmann equation with apparent valence and $V_h$ values of 1.14 and 17 mV for the native cell data and 1.57 and 120 mV for the heterologously expressed BK channel.
of the Boltzmann equation. It is apparent that the activation of BK channels in the parotid cells occurred at potentials approximately 120 mV more hyperpolarized than the same channels in the heterologous expression system. There also appeared to be a somewhat smaller voltage sensitivity of activation of the native channels as judged by the small decrease in the z parameter.

In several such experiments with 80 nM Ca^2+ we found a mean value for the V_h values of BK channels in native cells of 46.3 ± 1.6 mV (n = 8) and, about 113 mV more depolarized, 159 ± 2.0 mV (5) for the heterologously expressed channels. The effective valence, z, also differed between the native and expressed channels but not nearly as dramatically: mean values of 1.2 ± 0.06 and 1.8 ± 0.09, respectively. The very hyperpolarized activation range of mouse parotid BK channels was also found in human parotid acinar cells. The difficulty of obtaining human tissue limits the amount of data that can be obtained but in 5 cells from two subjects the mean V_h and z values were 61 ± 3.5 mV and 1.6 ± 0.11, respectively. This property was not confined to the parotid gland: acinar cells from the mouse submandibular gland (SMG) exhibited a V_h value of 37 ± 1.8 mV (9), even more hyperpolarized than parotid cells, and a z value of 1.79 ± 0.06 (9). The human parotid and mouse SMG data were obtained in the absence of crown ether (see Materials and Methods) but this is not likely to be a major issue at this low Ca^2+ concentration.

The large difference between the activation voltage ranges of native channels and those expressed in the CHO cells continued as the internal Ca^2+ was increased to 500 nM. Figure 2C shows raw currents recorded at this concentration from a native parotid cell (upper) and a cell in which the channels were heterologously expressed (note the different voltage ranges illustrated). Part D of the figure contains the relative conductance-voltage relation from these two cells. The large difference between the activation voltage ranges is apparent with mid point values of 17 and 120 mV for the channels in the native cell and those in the expression system. The steepness of these two relations also differed (z values of 1.14 and 1.57, respectively) and both appeared smaller than their counterparts at 80 nM Ca^2+. Pooled data from many cells confirmed these observations: mean V_h values for native cells was 16.5 ± 2.2 mV (12) and 124 ± 1.8 mV (4) and mean z values of 0.96 ± 0.047 (12) and 1.6 ± 0.11 (4), respectively.

Similar experiments were carried out over a range of Ca^2+ concentrations from near zero to 2 µM, which encompasses the physiological levels of Ca^2+ in these cells.11-14 The pooled results from these experiments are illustrated in Figure 3. The activation curves from BK channels in the native cells are shown with filled symbols and those from heterologously expressed channels are in open symbols. The large difference between the activation range of channels in these two cell systems is apparent. At all concentrations of Ca^2+, the activation range of the native channels occurred at substantially more hyperpolarized values than for expressed channels.

It is useful to compare the voltage dependence of BK channel activation in the nominal absence of Ca^2+ in order to remove complications from Ca^2+-dependent processes. At these “zero” Ca levels the mid-point of parotid cell BK channel activation and activation of the same channels expressed in CHO cells were 51.4 ± 2.0 mV (5) and 168 ± 3.2 mV (5), respectively. That is, the native channels activated at potentials almost 120 mV more hyperpolarized than the heterologously expressed channels. The effective valence, z, was much more similar: 1.24 ± 0.088 (5) and 1.42 ± 0.076 (5) for the native and expressed channel, respectively. These data are listed in Table 1 along with the range of published data from heterologously expressed mSlo and hSlo channels. These data show that, in the nominal absence of Ca^2+, the voltage activation of expressed parSlo BK channels was quite similar to the mSlo and hSlo variants and all of these are substantially more depolarized than the channels in native parotid cells.

**Figure 3.** Voltage- and Ca^2+ -dependence of mean BK channel relative conductance from native cells and from cells expressing parSlo. Data from 4–12 cells at each Ca^2+ level. Lines are fits of the standard Boltzmann equation.

**Table 1.** BK channel activation parameters in nominally zero Ca^2+.

| Channel                  | V_h (mV)       | z             |
|--------------------------|----------------|---------------|
| Native parotid BK         | 51.4 ± 2.0 (5) | 1.24 ± 0.088 (5) |
| Expressed parSlo          | 168 mV ± 3.2 mV | 1.42 ± 0.076 (5) |
| Expressed mSlo/hSlo       | 161–209 mV     | 0.95–1.3      |
| β1/β4 (+/+, +/-)          | 39.2 ± 5.1 (5) | 1.14 ± 0.11 (5) |
| β1/β4 (-/-, -/-)          | 44 ± 2.6 (6)   | 1.25 ± 0.089 (6) |

Shown are the midpoints (V_h) and effective valence (z) from fits of the Boltzmann equation to BK channel activation data. The range of values for the expressed mSlo and hSlo channels are examples from previously published studies.18-23
In many cells BK channels are heteromeric complexes with one of four types of accessory subunits, \( \beta_1 \) to \( \beta_4 \). The presence of these subunits alters the activation voltage range \(^{8,19,21,23-27} \) so it is possible that the hyperpolarized shift of the native channels could be due to the presence of one of these subunits. Parotid BK channels are not likely to express \( \beta_2 \) or \( \beta_3 \) subunits since these induce an inactivation in the channels not seen in the native channels. The lack of \( \beta_2 \) and \( \beta_3 \) transcripts in parotid cells is confirmed by PCR analysis of tissue from the parotid gland showing the presence of only \( \beta_1 \) and \( \beta_4 \).\(^1\) In general these subunits produce a depolarizing shift of BK channel activation in low \( \text{Ca}^{2+} \) so their presence should not be responsible for the hyperpolarized activation of native BK channels seen in low \( \text{Ca}^{2+} \). In order to verify this assertion, we generated mice with both the \( \beta_1 \) and \( \beta_4 \) genes ablated. We compared the BK channel activation in these \( \beta_1/\beta_4 \)-null mice to that in mice with the same genetic background but with both beta subunit genes intact. We found mean values of \( V_h \) and \( z \) of BK channels in parotid cells from the \( \beta_1/\beta_4 \)-null mice of \( 44 \pm 1.1 \) mV (6) and \( 1.25 \pm 0.04 \) (6), respectively. These values are not statistically different from their gene-matched control group and very similar to the values described above for BK channels from our standard animals. These data are included in Table 1.

**Figure 4A** provides a summary of the \( V_h \) values from native BK channels and the \( \text{parSlo} \) channels expressed in CHO cells over the range of \( \text{Ca}^{2+} \) examined. The large difference between the activation voltage range of BK channels in the native cells and in the heterologous expression system is apparent. These data seem to indicate that the \( \text{Ca}^{2+} \)-induced shift of \( V_h \) values of native channels was different than those in the expression system. This suggestion is confirmed by the data in **Figure 4B** where the shift in activation (from the “zero” \( \text{Ca}^{2+} \) level) produced by the indicated \( \text{Ca}^{2+} \) concentrations is shown for channels in native cells (■) and those in the expression system (○). At least over this range of \( \text{Ca}^{2+} \), these two sets of data can each be well-described by a standard single \( \text{Ca}^{2+} \) binding isotherm with apparent \( K_d \) values of 158 nM and 1,300 nM for native cell and CHO cell data, respectively. Solid and dashed red lines discussed in text.

**The functional basis for parotid BK channel activation.** It would be instructive to understand the mechanistic basis for the difference in the activation voltage range between the BK channels in native parotid acinar cells and these same channels in the heterologous expression system. The diagram below represents the current knowledge of the \( \text{Ca}^{2+} \)- and voltage-dependent behavior of BK channels derived from many studies over several years\(^{28-31} \) which has been given its most complete, quantitative description by Horrigan and Aldrich.\(^{17} \) In this Horrigan-Aldrich (HA) gating scheme, the voltage sensors of each of the four subunits undergoes a transition between a resting (R) and an activated (A) conformation governed by a voltage dependent equilibrium constant, \( J \). The actual channel opening is conferred by a transition from a closed (C) to an open (O) conformation with an equilibrium constant, \( L \), that is also voltage-dependent.
Ca\(^{2+}\)-binding and channel opening (C) and Ca\(^{2+}\)-binding and voltage sensor activation (E).

The mathematical description of the activation of the channels with this gating scheme is given by:

\[
P_o = \frac{L(1 + JD + KC + JKCD)^4}{L(1 + JD + KC + JKCD)^4 + (1 + J + K + JKE)^4}
\]

where

\[
J = \exp\left(\frac{z_J F (V_m - V_J)}{RT}\right)
\]

\[
K = \frac{[Ca^{2+}]}{K_d}
\]

\[
L = L_0 \exp\left(\frac{z_L F V_m}{RT}\right)
\]

\[V_J\] sets the voltage range of voltage sensor activation; it is the half-activation voltage for each voltage sensor in the absence of Ca\(^{2+}\). The parameter \(z_J\) controls the voltage sensitivity for activation of the voltage sensors and \(z_L\) controls the voltage sensitivity of the closed-open conformational change. \(K_d\) is the dissociation constant for a single calcium binding site.

In this scheme there are three parameters that strongly control the voltage-dependent activation of the channel in zero Ca\(^{2+}\): \(V_J\), \(L_0\), and \(D\). It has been shown\(^{17}\) that the voltage at which the channel activation time constant has its maximal value provides a rough estimate of the \(V_J\) value. Figure 5A shows the measured activation time constants for native BK channels in parotid acinar cells (■) and channels expressed in CHO cells (○). Similar to data obtained from expressed hSlo and mSlo channels,\(^{17-19}\) we found that the time constants for expressed parSlo channels has a maximum at very positive potentials—near 150 mV. In sharp contrast, the time constants for activation of BK channels in parotid cells peaks at much lower voltages—near zero mV. Thus, it is immediately clear that at least one of the mechanisms underlying the strongly hyperpolarized activation range of native BK is that the voltage sensors appear to operate at much more hyperpolarized potentials.

It is more difficult to determine the other two parameters (\(L_0\) and \(D\)) controlling the voltage range of channel activation.

The Ca\(^{2+}\) dependence of activation results from the binding of Ca\(^{2+}\) ions to a receptor (X) in each of the four subunits with equilibrium constant, \(K\). D, C and E are allosteric factors that control the synergy among these various gating components. Values of unity for these parameters represent no cooperativity; the larger the value the stronger the coupling. For example, the C-O equilibrium constant \(L\) increases D-fold for each voltage sensor that is activated. There are similar allosteric interactions between...
However, since it appears that the Ca\(^{2+}\) and voltage sensitivities of heterologously expressed parSlo channels are quite similar to hSlo and mSlo channels we can use previously determined values for these parameters (and for \(z_J\) and \(z_L\)) as starting points for our analysis of expressed parSlo channels. Figure 5B shows the voltage dependence of expressed parSlo channel open probability (relative conductance) in nominally zero Ca\(^{2+}\) (■) (from Fig. 3). The solid line is from the HA model with the indicated parameters. These parameter values are listed in Table 2 along with the range of those published for hSlo and mSlo channels. It is apparent that the values of these parameters from expressed parSlo channels were generally within the range of published values. One exception was the steepness of the closed-open conformational change (\(z_J\)) which was a bit larger in parSlo channels. Thus the HA model provided an excellent description of heterologously expressed parSlo channels with properties very similar to those of expressed hSlo and mSlo channels previously reported.

We then turned our attention to the determination of the HA model parameters for the native BK channels. The time constant data of Figure 5 indicates that the \(V_J\) parameter will be in the vicinity of 0 mV. Determining the \(L_0\) and \(z_L\) parameters is more difficult. One approach involves measuring gating charge movements.\(^{17,18}\) We were unable to detect gating currents in the native cells most likely because of the relatively few channels expressed in these cells. An alternate method to estimate these parameters is to measure the channel open probability at very negative potentials. At such voltages there is a very low channel open probability and the voltage sensors are all in their resting state. Under these conditions channel activation is described by:

\[
P_O = \frac{L}{1 + L}
\]

For \(L \ll 1\) (again at negative potentials where \(P_o\) is very small) this becomes:

\[
P_O = L = L_0\exp\left(\frac{z_L F V_m}{RT}\right)
\]

Thus, \(L_0\) and \(z_L\) can be estimated from measurements of channel open probability at very negative potentials where \(P_o\) is very small. Under these conditions the open channel probability will be a simple exponential function of \(V_m\) and so these will be linearly related on a semi-logarithmic plot. We determined the BK channel open probability of native BK channels at very negative potentials by measuring single BK channel activity in whole cell mode as described in Materials and Methods. We found an approximately linear relationship between \(P_o\) and \(V_m\) (open symbols in Fig. 5C) and a best fit to the data was obtained with \(L_0\) and \(z_L\) values of 0.02 and 1.78, respectively. It should be emphasized that these are only rough estimates for at least two reasons: (1) the inherent difficulty and potential technical problems in determining the open probability of single channels in whole cell, patch clamp mode and (2) the possibility that the conditions necessary for the analysis are not met. In particular, the voltage range used must be sufficiently negative to assure the voltage sensors are in the resting state. The data in Figure 5A shows that voltage sensor activation of native BK channels was substantially more hyperpolarized than of the expressed channels which raises concern about the rigorous validity of the assumption. Data could not be obtained at more negative voltages because of deleterious effects on the cells. Nevertheless, the values of \(L_0\) and \(z_L\) obtained with this method should still provide a reasonable approximation of the true values.

We used these estimates for the values of \(L_0\) and \(z_L\) and the estimate for \(V_J\), as described above, as starting points for determining a complete set of HA parameters that could describe the native channel open probability over the full range of experimentally accessible membrane potentials. The results of this analysis are illustrated in Figure 5C. The filled symbols are the open channel probability data (relative conductance) from Figure 3 for zero Ca\(^{2+}\). The open symbols represent the channel open probability obtained from single channel measurements in whole cell mode. The solid line is from the HA model with the indicated parameters and these are included in Table 2.

The results of these analyses show that the BK channels in native parotid acinar cells exhibited three significant differences from their heterologously expressed parSlo counterparts: (1) much larger \(L_0\) values which means the closed-open transition will occur at more negative potentials; (2) much more hyperpolarized \(V_J\) values, which means the voltage sensors operate at more negative values; and (3) a much smaller allosteric parameter \(D\). A small allosteric coupling would, by itself, cause channel activation to shift to more positive potentials because channel activation would rely more on the open-closed conformational change which occurs at potentials relatively positive to the voltage range for voltage sensor activation. Thus, the very hyperpolarized activation of native BK channels is due to the fact that both voltage sensor activation and the open-closed conformational change occur at more hyperpolarized potentials—in spite of a decreased coupling between these two processes.

The data that we have obtained on the Ca\(^{2+}\)-dependent shift of \(V_J\) (Fig. 4) should allow us to define the Ca\(^{2+}\)-dependent model parameters \(K_d\), \(C\) and \(E\). We expected to be able to describe the expressed parSlo data with parameters similar to the published

| Parameter | Expressed parSlo | Expressed mSlo/hSlo | Native BK |
|-----------|------------------|---------------------|-----------|
| \(L_0\)   | 0.82 x 10^-6     | 0.2–2.2 x 10^-6     | 0.001     |
| \(z_L\)   | 0.6              | 0.31–0.41           | 1         |
| \(V_J\)   | 148 mV           | 120–150 mV          | -20 mV    |
| \(z_J\)   | 0.45             | 0.5–0.58            | 1         |
| \(D\)     | 20               | 16.8–40             | 3.5       |
| \(K_d\)   | 8 µM             | 8.2–21.8 µM         | 0.8 µM    |
| \(C\)     | 11               | 8–14                | 2.3       |
| \(E\)     | 1.8              | 2.4–6               | 1         |

Shown are the HA parameters for the channels indicated. The various parameters are described in the text.
values. This was indeed the case as can be seen from the red line in Figure 4 that provides a good description of the data from the expressed channels. The $K_v$, $C$, and $E$ parameters for this description are listed in Table 2 and are quite similar to previously published values for hSlo and mSlo.

In contrast, we were not able to obtain a good fit to the data from the native cells with any set of Ca$^{2+}$-dependent parameters as long as the $K_v$ value was within the published range. The best version is the dashed red line in Figure 4B associated with the data from the native cells and with values for $K_v$ and allosteric parameters, $C$ and $E$, of 8 µM, 5, and 1.8, respectively. In order to provide an accurate description of the Ca$^{2+}$ dependence of the BK channels in native cells (solid red line in Fig. 4B) a 10-fold lower $K_v$ value of 0.8 µM was required along with $C$ and $E$ parameters of 2.3 and 1, respectively. These values are included in the summary in Table 2.

**Discussion**

**Parotid IK1 channel Ca$^{2+}$ sensitivity.** In this study we examined the Ca$^{2+}$ dependence of the activation of IK1 and BK channels in parotid acinar cells. Parotid IK1 channels were activated by Ca$^{2+}$ with an apparent $K_v$ value near 360 nM and a Hill coefficient of about 3. These values are similar to those for IK1 channels in other native cells and in heterologous expression systems. Thus, the Ca$^{2+}$ sensitivity of IK1 channel activation appears quite similar across species and cell types.

**Parotid BK channel Ca$^{2+}$ sensitivity.** In contrast to the global uniformity of the properties of IK1 channels in native cells and in expression systems, we found that the Ca$^{2+}$ and voltage sensitivity of BK channels in mouse parotid acinar cells were quite different than the analogous channels in other native cells (e.g., skeletal muscle, arterial smooth muscle and neurons) and expression systems. Specifically, in very low intracellular Ca$^{2+}$ the BK channels in parotid acinar cells activated at potentials more than 100 mV more hyperpolarized than BK channels in native excitable cells and the different Slo variants in heterologous expression systems. This property of BK channels is not restricted to mouse parotid acinar cells—human parotid and mouse submandibular cells have a similarly hyperpolarized voltage activation range (see Table 1). We found that BK channels in mouse distal colon crypt cells have that same property (unpublished observations). Cochlear inner hair cells also express BK channels$^{40}$ with very hyperpolarized activation in low Ca$^{2+}$ as do channels in a prostate cancer cell line.$^{37}$ This shared property suggests that it may be generally true that cells of epithelial origin all express native BK channels with a hyperpolarized activation voltage range.

We also investigated the voltage dependence of parotid BK channels with increased levels of Ca$^{2+}$. We found that the hyperpolarizing activation shift caused by increased Ca$^{2+}$ tended to saturate at values near 40 mV for Ca$^{2+}$ levels of a few µM whereas expressed parSlo (like mSlo and hSlo) shifted near 100 mV for these modest Ca$^{2+}$ levels and was not nearing saturation.

**Functional basis for the hyperpolarized voltage activation range of parotid BK channels.** In order to understand the origin of the differences between BK channels in parotid cells and in heterologous expression systems, we analyzed our results in the context of the Horrigan-Aldrich (HA) description of BK gating. There is a cooperative relationship among three processes in BK channel activation (see Results): a voltage dependent activation of the four voltage sensors, the voltage dependent opening of the channels, and the Ca$^{2+}$ binding to receptors on the four channel subunits. The hyperpolarized activation of parotid BK channels could result from enhanced activation of the voltage sensors (an increase in the HA J parameter), an enhanced likelihood of channel opening (L), or the allosteric parameter (D) that couples these two processes. Our analysis revealed that the parotid BK channels have a strongly reduced (6-fold) coupling between the voltage sensor activation and channel opening. This property on its own would result in a depolarized shift of BK activation but both the voltage sensor activation and channel opening process were very much shifted to more hyperpolarized potentials.

We found that the HA model could accurately describe the full range of the voltage dependence of parotid acinar cell BK channels and expressed parSlo channels at low Ca$^{2+}$. We also showed that this model could account for the shift of $V_v$ with increased Ca$^{2+}$ for both these channels—by increasing the apparent Ca$^{2+}$ affinity of the native channels. While the HA model accurately described the voltage steepness in nominally zero Ca$^{2+}$, it could not account for the observed small decrease in this parameter with increased Ca$^{2+}$ without an accompanying change in the parameters that control voltage steepness. In contrast, while the fits are not perfect, there are examples of the application of the HA model to expressed BK channels that can reasonably account for the small, Ca$^{2+}$-dependent increase in voltage steepness.$^{17-19,21}$

We do not know the origin of the inability of the HA model to accurately describe the voltage steepness of parotid BK channel activation over a range of Ca$^{2+}$ concentrations but there are examples of the same inability for BK channels expressed with a beta subunit.$^{18,19,21}$

Thus, compared with their expressed channel counterparts, the parotid BK channels seem to have (1) a very different voltage range for activation of the voltage sensors; (2) a very different voltage range for channel opening; (3) a large reduction in the coupling between voltage sensors and channel opening; (4) an increased Ca$^{2+}$ binding affinity; (5) some other, more minor, alteration that accounts for the inability of the HA model to accurately describe the Ca$^{2+}$-induced changes in the steepness of the voltage dependence of activation. The net result of these differences is BK channels that are seemingly optimized to operate in the physiological range of intracellular Ca$^{2+}$ in parotid acinar cells—without the need for these channels to be in microdomains of very high Ca$^{2+}$ levels. It would be instructive to determine if the functional basis for the low Ca$^{2+}$, hyperpolarized activation of BK channels in the other cell types is similar to that for the BK channels in native parotid cells.

**Molecular basis for the hyperpolarized activation of parotid BK channels.** We do not know the molecular basis for the hyperpolarized activation range of parotid BK channels but possibilities include interaction with auxiliary proteins, post-translational modifications, phosphorylation, oxidation-reduction effects, and
the actions of other modulatory agents. The category of auxiliary proteins could include the four known BK channel subunits. Of these, parotid tissue shows evidence of expression of only β1 and β4. All four subunits alter the relationship between voltage and Ca2+ activation but they actually make the BK channels less active at low Ca2+ levels and so would not likely underlie the high activity of parotid BK channels in low Ca2+. Nevertheless, we tested the BK channels in parotid acinar cells from animals with both the β1 and β4 genes ablated and these had the same hyperpolarized gating range in low Ca2+ as wild-type animals. Thus, the high activity of native parotid BK channels in low Ca2+ cannot be explained by the presence of one or both of these two β subunits. Note that upregulation of one or both of the other two β subunits can be ruled out since each of these subunits causes clear inactivation of the BK channel which is not seen in the parotid channels.

BK channels are subject to posttranslational modification, phosphorylation by serine-threonine kinases,49 tyrosine kinases,50 hormones,41 and the actions of redox agents42 and other small molecules.43 The effects of all these modifications and reagents result in altered activation of BK channels. However, these changes are quite modest compared to the 120 mV differences between native parotid BK channels and most other BK channels.

The modest effects of posttranslational modifications and modulation by kinases or small molecules would seem to make it unlikely that anything similar would be able to account for BK activation at relatively hyperpolarized potentials. While, as noted above, the known β subunits produce a shift of the voltage dependence of BK channel activation in the opposite direction needed, all but β3 can cause up to a 60 mV hyperpolarizing shift in elevated Ca2+.25,27 Thus, while we do not know the actual basis for the hyperpolarized activation of parotid (and other epithelial) BK channels, we speculate that it may be produced by association with a protein or other large molecule that has not yet been shown to modify BK channel gating.

**Parotid BK channel physiology.** The fluid secretion process requires a movement of Cl− ions through Ca2+-activated Cl channels (CaCCs) in the apical membrane of secretory acinar cells. Without the concomitant activation of K channels, secretion would come to an immediate halt as the increased Cl− conductance would drive the membrane potential to the Cl− equilibrium potential. The apparent Ca2+ affinity for CaCC activation is strongly voltage-dependent and at physiological voltages is near 300 nM.45,46 In this study we found a value of about 350 nM for the Ca2+ affinity of parotid IK1 channels at physiological voltages. We also found that a reasonable fraction of BK channels would be activated at a few hundred nM of Ca2+ even at physiological voltage levels. While the fraction of BK channels activated under these circumstances is not large, single BK channels have about 8 times the conductance of IK1 channels and about 50 times the conductance of the CaCCs,45,47,48 so even a relative few BK channels can exert a significant influence on the physiology of secretory cells. We showed in this study that the particular biophysical properties of salivary BK and IK1 channels endow them with the necessary characteristics to coordinate with CaCCs in order accomplish their physiological roles in fluid and electrolyte secretion.

**Materials and Methods**

**Ethical approval.** The procedures for animal handling, maintenance and surgery were approved by the University of Rochester Committee on Animal Resources. All animals used in this study were housed in a pathogen-free area at the University of Rochester. Mice were sacrificed by exsanguination following exposure to CO2.

Human parotid tissue was obtained from healthy subjects (30–70 years of age) undergoing surgical removal of a pleomorphic adenoma. Excess normal tissue surrounding the tumor is not used for diagnostic evaluation of the sample. This discarded tissue was collected immediately after surgical excision and transported to the laboratory in ice-cold physiological saline and acinar cells were isolated for electrophysiological recordings. These procedures were approved by the University of Rochester Institutional Review Board.

**Parotid acinar cell preparations.** Detailed descriptions of the procedures for producing single parotid acinar cells have been previously described. Briefly, surgically removed parotid glands were finely minced and digested with trypsin after which the cells were dissociated with Liberase and, finally, plated onto 5 mm diameter glass coverslips. All dissociation solutions were gassed continuously with 95% O2 + 5% CO2 and maintained at 37°C. Several strains of mice were used in this study. Most of the data were obtained from BlackSwiss x 129/SvJ hybrid animals. Some data were collected from mice in which genes encoding both the BK β1 and β4 subunits were ablated. These animals were the result of breeding β1(-/-) and β4(-/-) mice (each in a C57 background) in the laboratory of James E. Melvin of this institution.

**Heterologous expression.** The mouse parotid KCa1.1 variant (parSlo) was cotransfected (0.5 µg or 1 µg) into CHO-K1 cells using a Nucleofector machine (Amaxa Biosystems, MD) according to the manufacturer’s instructions or via Lipofectamine LTX (Invitrogen, CA) along with 0.25 µg of the fluorescent vector EYFPdcc315 for transfected cell identification. Cells were used between 24 and 48 h after transfection. The CHO-K1 cells were obtained from ATCC and grown in Ham’s F-12K medium + L-glutamine (ATTGC, VA) with 10% FBS and maintained in a CO2 incubator with 5% CO2.

**Electrophysiological recordings.** Membrane currents were acquired (at 20–22°C) using Axopatch 200B amplifiers and a Digidata 1320A digitizer (Axon Instruments, Foster City, CA) or a 12-bit analog/digital converter system of our own design. Membrane voltages were corrected for the measured junction potential between the external and internal solutions. Pipettes from quartz patch glass (Garner Glass Co., Claremont CA) were used and measured access resistance values were between 2.5 and 8.5 MΩ with a mean value of 5.7 ± 0.19 (SEM, n = 59). Owing to the large macroscopic currents in the native cells special attention
was paid to series resistance compensation. A minimum value of 90% was used with most values greater than 95%. The external solution contained (in mM): 150 Na-glutamate, 5 K-glutamate, 2 CaCl₂, 2 MgCl₂, 10 HEPES, pH 7.2. The internal solution contained 135 mM K-glutamate buffered to pH 7.2 with 10 mM HEPES and various free Ca²⁺ concentrations designed with MaxChelator Sliders (Chris Patton) (http://maxchelator.stanford.edu). The Ca²⁺ buffers used for the various free concentrations were: 10 mM BAPTA (80 nM); 2.5 mM EGTA/2.5 EDTA mM (500 nM); and 5 mM HEDTA (2 µM). A nominally 0 Ca²⁺ solution contained 10 mM BAPTA with no added Ca²⁺ with an estimated free Ca²⁺ level of 3 nM. All internal solutions contained 50 µmol/l crown ether to chelate potential Ba²⁺ contamination. The use of glutamate as the main anion in these solutions was used to reduce Cl⁻ channel currents in the native parotid cells to insignificant levels.

The external solutions used to isolate currents through BK channels in the parotid acinar cells included maurotoxin (Sigma) and TRAM-34 (Sigma) to eliminate currents through IK1 channels. The TRAM-34 concentration used was 1 µM from a 10 mM stock in DMSO. Maurotoxin was used at a final concentration of 50 nM in the presence of 0.01% BSA from a 50 µM stock solution in 15 mM Na₂HPO₄. Maurotoxin concentrations as high as 1 µM have a negligible effect on BK channel activity. This was done in the whole cell, patch clamp mode with an external solution consisting of (in mM): 135 K-glutamate, 2 CaCl₂, 2 MgCl₂, 10 HEPES, pH 7.2. In these experiments we measured the fractional single channel open time and, with the measured single channel current level, computed the conductance. The maximum BK conductance was obtained in the same cell via macroscopic current measurements and the relative conductance computed as the ratio of these.

IK1 channels are voltage- and time-independent and so we measured the currents through these channels from the instantaneous change in current (immediately after the settling of the capacity transient) at each test potential before significant activation of the time-dependent BK channel. Negligible IK1 channel currents are activated at Ca²⁺ concentrations below 100 nM. IK1 channels are activated by an increase in the open channel probability by the chemical compound DCEBIO with a Kₘ value of about 840 nM. The DCEBIO activation of IK1 channels is independent of intracellular Ca²⁺ levels above 100 nM. Thus, in our experiments we determined the relative level of IK1 activation from currents recorded at a given level of intracellular Ca²⁺ and the current level induced by 20 µM DCEBIO.

**Data analysis.** BK channel steady state currents (Iₛₛ) were used to compute conductance-voltage (G-V) relation: $G = \frac{G_{\text{max}}}{1 + \exp^{-z(V_m-V_h)/R\beta T}}$, where $G_{\text{max}}$ is the maximum conductance, $z$ is the effective charge, $V_m$ is the membrane voltage and $V_h$ is the potassium equilibrium potential. Relative channel activation was determined as the relative BK conductance. The G-V relations in low Ca²⁺ did not reach a saturating level at reasonably accessible voltages so in these cases the fit of a Boltzmann function was used to estimate the maximal level.
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