Gating of Recombinant Small-Conductance Ca-activated K⁺ Channels by Calcium

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ABSTRACT Small-conductance Ca-activated K⁺ channels play an important role in modulating excitability in many cell types. These channels are activated by submicromolar concentrations of intracellular Ca²⁺, but little is known about the gating kinetics upon activation by Ca²⁺. In this study, single channel currents were recorded from Xenopus oocytes expressing the apamin-sensitive clone rSK2. Channel activity was detectable in 0.2 μM Ca²⁺ and was maximal above 2 μM Ca²⁺. Analysis of stationary currents revealed two open times and three closed times, with only the longest closed time being Ca dependent, decreasing with increasing Ca²⁺ concentrations. In addition, elevated Ca²⁺ concentrations resulted in a larger percentage of long openings and short closures. Membrane voltage did not have significant effects on either open or closed times. The open probability was ~0.6 in 1 μM free Ca²⁺. A lower open probability of ~0.05 in 1 μM Ca²⁺ was also observed, and channels switched spontaneously between behaviors. The occurrence of these switches and the amount of time channels spent displaying high open probability behavior was Ca²⁺ dependent. The two behaviors shared many features including the open times and the short and intermediate closed times, but the low open probability behavior was characterized by a different, long Ca²⁺-dependent closed time in the range of hundreds of milliseconds to seconds. Small-conductance Ca-activated K⁺ channel gating was modeled by a gating scheme consisting of four closed and two open states. This model yielded a close representation of the single channel data and predicted a macroscopic activation time course similar to that observed upon fast application of Ca²⁺ to excised inside-out patches.

KEY WORDS: potassium channel • calcium-activated • single channel • afterhyperpolarization • gating

INTRODUCTION Intracellular Ca²⁺ is an important second messenger in most cell types. One pathway by which Ca²⁺ controls cellular functions is regulating the K⁺ efflux through the cell membrane. Ca²⁺-activated K⁺ channels have been demonstrated in many tissues and have been classified into large, intermediate, and small conductance Ca-activated K⁺ channels (BK, IK, and SK channels, respectively).¹ The three classes differ with respect to their single channel conductance, Ca²⁺ sensitivity, voltage dependence, and pharmacology (Blatz and Magleby, 1987; Latorre et al., 1989; Ishii et al., 1997). While BK channels have been studied extensively, the biophysical properties of SK and IK channels are less well characterized.

SK channels were first identified in cultured rat skeletal muscle, where they appeared to open independent

¹Abbreviations used in this paper: AHP, afterhyperpolarization; BK (and SK) channels, large (and small) conductance Ca-activated K⁺ channels.
The cloning of SK channels has revealed the existence of at least three members of a family (SK1–3). The amino acid sequence of the three subtypes was highly conserved, and hydrophobicity analysis predicted a secondary structure with six membrane-spanning regions. Further, two subtypes examined, rSK2 and hSK1, displayed a calcium dependence and single channel conductance similar to that observed for native channels (Köhler et al., 1996). As seen with native SK channels, differences in apamin sensitivity were observed for the cloned channels. For example, rSK2 was blocked by picomolar concentrations of apamin, whereas hSK1 was apamin insensitive (Köhler et al., 1996). Northern blot analysis showed the mRNA for rSK2 to be present in adrenal gland (Köhler et al., 1996), in agreement with the observation of apamin-sensitive SK channels in adrenal chromaffin cells (Park, 1994). In situ hybridization revealed that the mRNA encoding SK1 was expressed in rat hippocampus (Köhler et al., 1996).

SK channels underlie the prolonged afterhyperpolarization (slow AHP) following trains of action potentials in many neurons (Sah, 1996). The slow AHP differs between tissues in its time course and pharmacology. For example, in bullfrog sympathetic neurons (Pennefather et al., 1985; Goh and Pennefather, 1987), the slow AHP decays within several hundred milliseconds and is blocked by apamin. In contrast, the slow AHP recorded from hippocampal pyramidal neurons follows a slower time course, decaying over several seconds, and is insensitive to apamin (Lancaster and Adams, 1986; Storm, 1987). Both apamin-sensitive and -insensitive AHPs are present in rat and guinea pig vagal neurons (Sah and McLachlan, 1991; Sah, 1993) and cat cortical neurons (Schwindt et al., 1992). Apamin-sensitive and -insensitive AHPs from different tissues have properties suitable for a role in modulating the frequency of action potential firing (Larg and Ritchie, 1990; Neely and Lingle, 1992; Park, 1994; Yarom et al., 1985). This phenomenon has been studied in detail in hippocampal pyramidal neurons, where it is referred to as spike frequency adaptation (Madison and Nicoll, 1984).

In this study, the single channel properties of rSK2 and their dependence on voltage and calcium was examined. Two types of SK channel activity with very different open probabilities were observed, with the two behaviors sharing many characteristics. They were described by two open and three closed times. The open times and the short and intermediate closed times were the same for both types of channel activity, whereas the long closed time differed ~10-fold. Only the long closed time was Ca dependent. Although relative contributions of open and closed times differed between the two types of channel activity, elevated intracellular Ca$^{2+}$ favored longer openings and short closures in both behaviors.

**Methods**

**Channel Expression**

All experiments used the apamin-sensitive clone rSK2 from rat brain cDNA expressed in *Xenopus* oocytes. *Xenopus* frogs were handled in accordance with the institutional guidelines and underwent no more than two surgeries separated by at least 3 wk. Standard techniques were practiced during in vitro mRNA synthesis, and oocyte handling and injection (Adelman et al., 1992). Oocytes were studied 15–48 h after injection with 2 ng of mRNA.

**Solutions**

During recording, oocytes were bathed in 116 mM K-gluconate, 4 mM KCl, 10 mM HEPES, 5 mM EGTA, adjusted to pH 7.2 with KOH. To yield the reported concentration of free Ca$^{2+}$, CaCl$_2$ was added as calculated using published stability constants (Maret and Smith, 1974). For Ca$^{2+}$ concentrations >1 µM, EGTA was omitted from the bathing solution and gluconate was used to buffer free Ca$^{2+}$, assuming a stability constant for calcium gluconate of 15.9 M$^{-1}$ (Dawson et al., 1969). Electrodes were filled with a solution containing 116 mM K-gluconate, 4 mM KCl, 10 mM HEPES, pH 7.2.

**Electrophysiology**

All recordings were performed on excised inside-out patches (Hamill et al., 1981) using thick-walled quartz electrodes (13–15 MΩ). Membrane patches were voltage clamped with an Axopatch 200 amplifier using a CV 201A headstage (Axon Instruments, Foster City, CA). Continuous recordings were low pass filtered at 1 kHz with an eight pole Bessel filter (Frequency Devices Corp.), logarithmically binned with 20 bins per decade, and fitted by the sum of exponential probability density functions using the maximum-likelihood method. A correction was made for the rise time of the filter (Colquhoun and Sigworth, 1983) and all events were processed at room temperature. Except for Figs. 2 and 3, a holding potential of ~80 mV was used; however, a variable offset up to 20 mV occurred upon patch excision and was not corrected.

**Data Analysis**

Single channel recordings were analyzed with MacTac (Bruxton Corp., Seattle, WA) using the “50% threshold” technique to estimate event amplitudes and duration, and each transition was visually inspected before being accepted. Open- and closed-duration histograms were constructed using MacTacfit (Bruxton Corp.), logarithmically binned with 20 bins per decade, and fitted by the sum of exponential probability density functions using the maximum-likelihood method. A correction was made for the rise time of the filter (Colquhoun and Sigworth, 1983) and all bins were used for fitting. The number of statistically significant components was determined using the method of maximum likelihood ratios (Horn and Lange, 1985), which assumes a chi-squared distribution for 2- ln (likelihood 1/likelihood 2). Missed events were not corrected.

Open probability was calculated using a computer program (ReadEvents1.37; Dr. Scott Eliasof, Vellum Institute) to divide the sum of all open times by the total time of recording. All patches included in the analysis, except one, had periods of high channel open probability lasting long enough to establish that only a single channel was present in the patch. No double-level openings were seen in the patch that had only low channel open probability, and closed times were similar to the ones observed in other single-channel patches.

Single channel kinetics were simulated using CSIM 2.0 (Axon Instruments). Simulations were run at the same output rate as
the experimental sampling rate, 0.15 pA RMS of pseudo-random Gaussian noise was added to the data, and a Gaussian low pass filter was applied at 1 kHz. Resulting simulated data files were analyzed in the same fashion as channel recordings. Initial estimates of rate constants were obtained from the life times and relative amplitudes of each exponential component in the duration histograms, and from the channel open probability. The simulated traces and duration histograms were compared with the experimental data, and rate constants were adjusted to obtain good agreement. For the model containing three open and four closed states presented in Figs. 10 and 11, 46 estimates of rate constants were examined. Macroscopic currents were simulated using SCoP (Simulation Resources, Inc., Berrien Springs, MI). Values were expressed as mean ± SD and P values were derived from unpaired, two-tailed Student’s t tests. Results were considered significantly different at P < 0.01.

R E S U L T S

rSK2 Channels Were Activated by Intracellular Calcium

Macroscopic currents recorded from oocytes expressing rSK2 were Ca dependent (Köhler et al., 1996). At the single channel level, this dependence manifested itself as an increase in open probability as a function of increased intracellular Ca\(^{2+}\) concentration. Fig. 1 A shows traces recorded from an excised inside-out patch containing a single rSK2 channel. In these traces, openings became more frequent as the Ca\(^{2+}\) concentration was increased from 0.6 to 5 μM. In four patches, the average open probability was 0.42 ± 0.12 in 0.6 μM Ca\(^{2+}\) and 0.74 ± 0.16 in 1 μM Ca\(^{2+}\). Fig. 1 B shows a plot of the open probability as a function of Ca\(^{2+}\) concentration for the patch in A. Fitting of the data to the Hill equation yielded an EC\(_{50}\) of 0.74 μM and a Hill coefficient of 2.2, mean values from three patches were 0.62 ± 0.14 μM and 3.2 ± 1.0, respectively. The EC\(_{50}\) was the same as that observed using macropatches (0.63 ± 0.23, P > 0.1). The Hill coefficient was somewhat reduced from the value of 4.8 ± 1.5 seen in macropatches (Köhler et al., 1996), although the difference was not statistically significant (P > 0.01, see discussion).

rSK2 Channels Were Voltage Independent

To examine the voltage dependence of rSK2 single channel kinetics, inside-out patches were held at membrane potentials from -100 to +100 mV. Analysis of the current amplitudes as a function of voltage yielded

**Figure 1.** Calcium dependence of SK channel activity. (A) Recordings from an inside-out patch in different calcium concentrations at a holding potential of -80 mV. Traces show 2-s periods of the single channel recordings used to calculate the open probabilities in B. Channel openings are shown as downward deflections and calculated free intracellular [Ca\(^{2+}\)] is noted on the left. Channel current was slightly smaller in the trace recorded in 5 μM Ca\(^{2+}\) due to an off-set of ~11 mV that occurred upon switching to the EGTA-free solution (see METHODS). (B) Open probability as a function of intracellular calcium concentration. 30–60-s steady state recordings in 0.6, 0.8, 1.0, and 5.0 μM Ca\(^{2+}\) were used to calculate the open probability. Least squares fitting of the data to the Hill equation of the form \(P_o = P_{o,max} \cdot [\text{Ca}^{2+}]^n/(EC_{50}^n + [\text{Ca}^{2+}]^n)\) (where \(P_{o,max}\) is the maximal open probability in saturating [Ca\(^{2+}\)], and other terms have their usual meanings) yielded \(P_{o,max} = 81\%\), EC\(_{50}\) = 0.74 μM, and \(n = 2.2\).
a slope conductance of 10.3 ± 1.3 pS in three single channel patches (data not shown), similar to the value previously reported (Köhler et al., 1996). Traces recorded from the patch in Fig. 1 at −60 and −100 mV were shown together with the corresponding open-time distributions (left) and closed-time distributions (right) in Fig. 2. Traces recorded at the two membrane potentials showed no obvious differences in channel activity. Both traces contained short and long openings, and open-time distributions were best fit by the sum of two exponentials with time constants of ~1 and 11 ms. Independent of membrane potential, three exponentials were needed to adequately fit closed-time distributions. Time constants were similar at both membrane potentials and were ~0.7, 4.5, and 30 ms.

The voltage dependence of the single channel behavior is summarized in Fig. 3. As discussed in detail below, only some of the parameters describing SK channel kinetics were Ca\textsuperscript{2+} dependent. Since the three patches examined were exposed to different Ca\textsuperscript{2+} concentrations, the Ca\textsuperscript{2+}-dependent parameters were plotted individually for each patch, whereas Ca\textsuperscript{2+}-independent parameters were shown as mean values. Patch membrane voltage did not significantly affect channel open probability (Fig. 3 A). As seen in Fig. 2, open-time histograms were best fit by the sum of two exponentials. The mean time constants were plotted as a function of voltage in Fig. 3 B. Voltage did not significantly affect either open-time constant. In addition, the relative contribution of each time constant to the open-time histogram was similar at all voltages examined (Fig. 3 C).

Three exponentials were needed to fit the closed-time histograms in two patches, whereas two exponentials were adequate for a third patch (see Fig. 3, legend). The short and intermediate closed times, present in all three patches, were shown as mean values in Fig. 3 D, and the long closed time was plotted for the two patches individually in Fig. 3 E. As was the case for the open times, voltage did not significantly affect the three closed-time constants, as illustrated by the exponential fits (Fig. 3, D and E, solid lines). Finally, the relative amplitudes of the short (Fig. 3 F, closed symbols) and intermediate (Fig. 3 F, open symbols) closed-time components were also insensitive to voltage. Because membrane potential did not affect any of the open and closed times, a single holding potential of −80 mV (plus offset, see METHODS) was used during the remainder of this study.

Figure 2. SK channel activity at different holding potentials. 1-s traces recorded from the patch in Fig. 1 exposed to 1.0 μM Ca\textsuperscript{2+} at membrane potentials of −60 and −100 mV (A and B, respectively). Channel openings are shown as downward deflections. Open-time (left) and closed-time (right) histograms are shown below each trace. Fitting a sum of two exponentials to the open-time distributions yielded at −60 mV: 1.4 ms, 40%; 10.6 ms, 60%; and at −100 mV: 1.2 ms, 40%; 11.0 ms, 60%. Fitting a sum of three exponentials to the closed-time distributions yielded at −60 mV: 0.74 ms, 57%; 5.2 ms, 25%; 36 ms, 18%; and at −100 mV: 0.65 ms, 58%; 3.6 ms, 28%; 26.7 ms, 14%. Relative amplitudes were obtained from the fitted exponential distributions.
Ca\textsuperscript{2+} Affected the Longest Closed Time as well as Open- and Closed-Time Distributions

Fig. 4 shows 2-s segments of recordings from a single channel patch exposed to different concentrations of intracellular Ca\textsuperscript{2+}. As seen in Fig. 1, the open probability increased with higher Ca\textsuperscript{2+} concentration. The change in open probability was reflected in both the open- and closed-time distributions. In all three concentrations of Ca\textsuperscript{2+}, open-time histograms (Fig. 4, left) showed two components with lifetimes independent of Ca\textsuperscript{2+}. However, an increased concentration of intracellular Ca\textsuperscript{2+} resulted in a larger fraction of long openings. For the recordings in Fig. 4, 49\% of all openings in 0.4 \textmu M intracellular Ca\textsuperscript{2+} contributed to the long open-time constant compared with 66\% in 1 \textmu M Ca\textsuperscript{2+}. Independent of Ca\textsuperscript{2+} concentration, closed-time histograms showed three components (Fig. 4, right). The short and intermediate closed times were unaffected by a change in Ca\textsuperscript{2+} concentration (see Fig. 5). In contrast, the long closed time increased from 20 ms in 1 \textmu M to 60 ms in 0.4 \textmu M free Ca\textsuperscript{2+}. Ca\textsuperscript{2+} also affected the relative amplitude of the closed-time components, shifting the distribution towards the shorter closed times. For the patch in Fig. 4, 71\% of all closures in 1.0 \textmu M Ca\textsuperscript{2+} were of the short time-constant compared with 63\% in 0.4 \textmu M.

The Ca\textsuperscript{2+} dependence of the kinetic parameters is summarized in Fig. 5. Fig. 5, A and B show the mean open and closed times from five single channel patches.
as a function of Ca\textsuperscript{2+} concentration. Neither the open times nor the short and intermediate closed time changed notably over the range of 0.4 to 30 \(\mu\text{M}\) Ca\textsuperscript{2+}. In contrast, the long closed time was clearly Ca\textsuperscript{2+} dependent and linear regression analysis yielded a slope of \(-66\ \text{ms \mu M}^{-1}\) (Fig. 5 C). In addition, the relative contributions of open- and closed-time constants to the respective lifetime histograms were Ca\textsuperscript{2+} dependent. As shown in Fig. 5 D, the average relative amplitude of the fast open-time component decreased approximately fourfold when the Ca\textsuperscript{2+} concentration was increased from 0.4 to 30 \(\mu\text{M}\). Concurrent with the increasing relative contribution of long openings, elevated Ca\textsuperscript{2+} concentrations increased the occurrence of closures of the short time constant (Fig. 5 E) and decreased the number of closures of the long time constant (Fig. 5 F). In 5 and 30 \(\mu\text{M}\) Ca\textsuperscript{2+}, two exponentials corresponding to the short and intermediate closed-time constants were sufficient to describe closed-time histograms.

**SK Channel Behavior Changed Spontaneously**

The traces in Fig. 6 A represent part of a continuous recording obtained from an inside-out patch bathed in 1 \(\mu\text{M}\) Ca\textsuperscript{2+}. Approximately 13 s after patch excision (Fig. 6 A, arrow), the open probability of the channel

![Figure 4](image-url)

**Figure 4.** SK channel activity in different concentrations of Ca\textsuperscript{2+}. Data shown are from a single channel patch voltage clamped at \(-80\ \text{mV}\). Scaling is the same for all traces and open probabilities are noted above each trace. Open-duration histograms are displayed on the left and closed-duration histograms on the right underneath the corresponding 2s traces. Solid lines and time constants represent maximum likelihood fits to the sum of two or three exponential components shown for each component separately as well as for the sum. Relative amplitudes of open-duration components were in 1.0 \(\mu\text{M}\) Ca\textsuperscript{2+}: 1.1 ms, 34%; 11.9 ms, 66%; in 0.6 \(\mu\text{M}\) Ca\textsuperscript{2+}: 1.0 ms, 43%; 10.1 ms, 57%; in 0.4 \(\mu\text{M}\) Ca\textsuperscript{2+}: 1.1 ms, 51%; 13.5 ms, 49%. Relative amplitudes of closed-duration components were in 1.0 \(\mu\text{M}\) Ca\textsuperscript{2+}: 0.8 ms, 71%; 4.2 ms, 19%; 20.0 ms, 10%; in 0.6 \(\mu\text{M}\) Ca\textsuperscript{2+}: 0.7 ms, 67%; 4.0 ms, 23%; 32.6 ms, 10%; in 0.4 \(\mu\text{M}\) Ca\textsuperscript{2+}: 0.8 ms, 63%; 4.8 ms, 25%; 60.2 ms, 12%.
decreased spontaneously. In this example, the decline of activity appeared to be somewhat gradual. However, in other examples, channel behavior changed more abruptly. Open probability remained low for \( \sim 47 \) s before it reverted back to the starting level (Fig. 6 A, arrow). Fig. 6 B shows the open probability during 1-s intervals plotted as a function of time after patch excision. During the segment of lower activity, the open probability was 0.03 compared with 0.81 during the combined segments of high activity.

During the period of low channel activity, short and long openings were seen similar to what was observed during periods of high activity, but openings or clusters of openings were separated by longer closures. The similarity became more obvious when openings during the high and low open probability periods were analyzed separately. Open times of 0.9 and 9.4 ms vs. 0.9 and 8.9 ms were obtained for high and low channel activity, respectively, with more of the openings being of the long time constant during high open probability (74%, compared with 35% during low open probability). Closed-time histograms representing the high open probability behavior showed three components of 0.6 (72%), 4.2 (23%), and 34 (5%) ms. Three closed-time constants were also seen during the low open probability period, and the short and intermediate components had similar dwell times of 0.6 (37%) and 5.1 (32%) ms, respectively. However, the third, long closed-time component had a dwell time of 320 ms and accounted for 31% of closures (data not shown, but see Fig. 8).

Periods of high and low open probability behavior were seen in 8 of 13 patches. Switches occurred either within continuously acquired recordings (16 examples in six patches) or during times between recordings.

Figure 5. Calcium dependence of channel open and closed durations. Data were from five excised patches, each containing a single channel displaying high open probability behavior (see text). Except in C, lines were drawn to illustrate the Ca\(^{2+}\) independence of the kinetic parameters and are not intended to have physical meaning. (A) Average short and long open times were independent of intracellular Ca\(^{2+}\) concentration. (B) Average short and intermediate closed-times were independent of Ca\(^{2+}\) concentration. (C) Ca\(^{2+}\) dependence of the average long closed times. Linear regression analysis of the means yielded a slope of \(-96 \text{ ms} \mu\text{M}^{-1}\). (D) The relative amplitude of the short open-time component decreased as a function of Ca\(^{2+}\) concentration. (E) The relative amplitude of the short closed-time component increased with increasing Ca\(^{2+}\) concentration. (F) Increasing Ca\(^{2+}\) concentrations lead to a decrease in the relative amplitude of the long closed-time component.
when the intracellular Ca$^{2+}$ concentration was changed (nine examples in five patches). The latter switches were always to high open probability when the Ca$^{2+}$ concentration was raised and to low open probability when Ca$^{2+}$ was decreased. Spontaneous switches in constant Ca$^{2+}$ were most often seen in Ca$^{2+}$ concentrations close to the EC$_{50}$. No switches were seen at the lowest (0.2 µM) or the highest (30 µM) Ca$^{2+}$ concentration examined, and in the two patches exposed to 0.2 µM free Ca$^{2+}$ for a total of 565 s, channels exhibited only low open probability. In contrast, channels showed only high open probability when exposed to 30 µM Ca$^{2+}$ (three patches) and out of two patches exposed to 5 µM Ca$^{2+}$ for 80 and 100 s, only one showed a very brief (4-s) period of low open probability. These observations suggested that the preference for each type of channel activity was Ca$^{2+}$ dependent. To illustrate this point, Fig. 7 A shows the total time of recording (all patches combined) exhibiting the two types of behavior for each Ca$^{2+}$ concentration tested. In the inset, the time during which high open probability was observed was normalized to the total recording time in each concentration of Ca$^{2+}$. Fitting of the data to the Hill equation yielded an EC$_{50}$ of 0.52 µM and a Hill coefficient of 4.1. Since, on the time scale of a typical recording, switches occurred infrequently, it was not possible to directly measure the overall open probability including both behaviors. However, since the low open probability (0.04 in 1 µM Ca$^{2+}$) was essentially negligible compared with the high open probability, an “effective open probability” was calculated by multiplying each single channel open probability by the relative time channels spent in high open probability behavior ($P_{o(high)} \cdot f(high)$) (Fig. 7 A, inset). Lines represent fits to the Hill equation, yielding an EC$_{50}$ of 0.43 µM and a Hill coefficient of 3.3 for the single channel open probability, and an EC$_{50}$ of 0.63 µM and a Hill coefficient of 3.6 for the “effective open probability.”
open probability" (Fig. 7 B, □) was calculated by multiplying the single channel open probability from a recording of high open probability behavior (Fig. 7 B, ●; EC$_{50}$ = 0.43 μM, Hill coefficient = 3.3) by the relative amount of time channels spent exhibiting high open probability behavior (Fig. 7 A, inset). Fitting the effective open probability to the Hill equation predicted a macroscopic EC$_{50}$ of 0.63 μM and Hill coefficient of 3.6 (see discussion).

A Ca$^{2+}$-dependent Time Constant of Hundreds of Milliseconds to Several Seconds Characterized Periods of Low Open Probability

Most patches exhibited periods of low channel open probability (see above) lasting from a few seconds to >15 min. Channels were observed to switch between behaviors (see above), and no double-level openings were seen. In addition, the single channel conductance and the two open-time constants were shared between both types of channel activity. These findings suggested that the channels underlying both types of activity were the same.

Fig. 8 shows steady state recordings from a patch containing a channel that displayed predominantly low open probability activity (0.05 in 1.0 μM calcium). The same patch showed high open probability for almost 3 min, revealing that only a single channel was present. Channel activity during low open probability periods was Ca sensitive and open probability increased as a function of Ca$^{2+}$ concentration by a factor similar to that in high open probability behavior (Fig. 8). During low open probability periods, SK channel activity exhibited similar open state kinetics as observed during high open probability. Independent of the Ca$^{2+}$ concentration, open-duration histograms were best fit by the sum of two exponentials (Fig. 8). Average time constants in seven patches were 0.93 ± 0.23 and 9.9 ± 4.4 ms. These open times were not statistically different from those observed during high open probability activity (1.1 ±
0.3 and 10.6 ± 2.9 ms, eight patches, \( P > 0.05 \). However, an obvious difference between the two types of SK channel activity was that short-duration openings predominated during low open probability behavior. For example, 71% of the openings observed in 1.0 \( \mu M \) free \( \text{Ca}^{2+} \) contributed to the short open-time constant during low open probability behavior (Fig. 8), while only 34% of openings were of short duration during high open probability activity (Fig. 4). Despite this difference, the relative amplitude of the short open-time component decreased with increasing \( \text{Ca}^{2+} \) concentration in both types of channel behavior. For example, in the patch shown in Fig. 8, 29% of openings contributed to the long open-time constant in 1.0 \( \mu M \) free \( \text{Ca}^{2+} \), compared with 17% in 0.4 \( \mu M \).

SK channel closed state kinetics during low open probability activity exhibited three exponential components (Fig. 8). The average short and intermediate closed-time constants of 0.9 ± 0.4 and 5.9 ± 1.5 ms (seven patches) were not statistically different from those observed during high open probability activity (0.7 ± 0.2 and 4.6 ± 1.3 ms, eight patches, \( P > 0.01 \)), but accounted for a smaller fraction of closures (60 ± 12% vs. 92 ± 6% in 1.0 \( \mu M \) \( \text{Ca}^{2+} \), three and four patches, \( P < 0.01 \)). In contrast, the third closed-time constant was significantly longer, measuring 295 ± 26 ms in 1.0 \( \mu M \) \( \text{Ca}^{2+} \) during low open probability, compared with 25.9 ± 5.8 ms during high open probability (five and three patches, respectively, \( P < 0.01 \)). As observed for high open probability behavior, the longest closed time was \( \text{Ca}^{2+} \) dependent and decreased from 1,290 ± 320 ms to 295 ± 26 ms (\( P < 0.01 \), three patches) when the free \( \text{Ca}^{2+} \) concentration was increased from 0.4 to 1.0 \( \mu M \) (see Fig. 9 C). During both types of channel behavior, higher \( \text{Ca}^{2+} \) concentrations shifted the distributions toward the shorter closed-time constants. For example, in the patch in Fig. 8, the longest closed-time constant accounted for 92% of closures in 1.0 \( \mu M \) and 43% in 0.4 \( \mu M \) \( \text{Ca}^{2+} \).

The \( \text{Ca}^{2+} \) dependence of the kinetic parameters describing low open probability behavior was summarized in Fig. 9. As seen during high open probability behavior (compare Fig. 4), neither the open times (Fig. 9 A) nor the short and intermediate closed times (Fig. 9 B) were \( \text{Ca}^{2+} \) dependent. The long closed time decreased with increasing \( \text{Ca}^{2+} \) concentration, and linear regression analysis yielded a slope of \(-2.5\) s \( \mu M^{-1} \) (Fig. 9 C). The relative contributions to lifetime histograms of the short open time (Fig. 9 D) and the short and long closed-time components (Fig. 9, E and F, respectively) differed from those observed during high open probability behavior, but they showed the same trend with increasing \( \text{Ca}^{2+} \) concentration; i.e., the number of short openings and long closures decreased, whereas the number of short closures increased.

**Discussion**

This report describes the effects of membrane potential and intracellular \( \text{Ca}^{2+} \) on the single channel kinetics of a recombinant apamin-sensitive SK channel. Previously, little was known about the behavior of single SK channels, with the only available kinetic data being obtained from native apamin-insensitive channels and currents (Lancaster et al., 1991; Valiante et al., 1997; Sah and Isaacson, 1995; Sah, 1995). As might be expected from the high degree of homology between apamin-sensitive and -insensitive recombinant subtypes (66% amino acid identity between \( \text{rSK2} \) and \( \text{hSK1} \); Köhler et al., 1996), results from these studies were similar to the results reported here for cloned apamin-sensitive SK channels. Apamin-insensitive SK channels recorded from cultured rat hippocampal neurons possessed open times that were fit with a single exponential time constant of 7.8 ms and closed times that were fit with two exponentials with time constants of 0.49 and 1.49 ms. These channels exhibited an open probability of \(~0.5\) in 1 \( \mu M \) \( \text{Ca}^{2+} \) (Lancaster et al., 1991). Analyses of current fluctuations during the slow AHP in dentate granule neurons yielded similar results. For example, the mean channel open time was 6.9 ms throughout the time course of the slow AHP, whereas the decay of the slow AHP was accompanied by an increase in the mean closed time (Valiante et al., 1997). Assuming that the decay of the slow AHP resulted from the clearance of \( \text{Ca}^{2+} \) from the vicinity of the channels, this finding agrees with our observation that only the longest closed time was \( \text{Ca} \) dependent. Somewhat shorter open times of 1.7 and 3.8 ms were measured using fluctuation analysis of \( I_{\text{AHF}} \) in hippocampal pyramidal neurons (Sah and Isaacson, 1995) and vagal motoneurons (Sah, 1995), respectively.

**SK Channel Activation Could Be Modeled as a Markov Process**

The activation of SK channels may be described as a time-homogeneous Markov chain. Single rSK2 channels recorded from oocytes showed open-time distributions that were best fit by the sum of two exponentials, whereas three exponentials were necessary to fit closed-time distributions. The high open probability behavior could be modeled using a gating scheme with two open and four closed states as illustrated in Fig. 10, where the forward transition rates between closed states were dependent on intracellular \( \text{Ca}^{2+} \). Strictly sequential gating schemes with two open states that were either connected or separated by closed states did not fit the data. Several gating schemes containing closed loops were also examined and did not yield satisfactory results. A scheme similar to the one presented in Fig. 10 but consisting of two open and three closed states yielded a rea-
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A reasonable representation of the data, but only if rate constants were allowed to have a nonlinear dependence on Ca\(^{2+}\) concentration. Since Ca\(^{2+}\)-dependent transitions were assumed to reflect simple bimolecular reactions, their rates should vary linearly with Ca\(^{2+}\) concentration. This was achieved by adding a fourth closed state. Due to the similarity of two of the predicted time constants, the gating scheme in Fig. 10 yielded closed-time histograms with three resolvable components rather than the theoretical four components. The rate constants were adjusted such that the predicted lifetimes and relative amplitudes were similar to what was observed experimentally for a representative patch exposed to two Ca\(^{2+}\) concentrations (compare Figs. 4 and 10). As seen in single channel recordings, channel open times and the short and intermediate closed times were unchanged when channel activity was simulated with rate constants for 0.4 and 1.0 \(\mu\)M free Ca\(^{2+}\), respectively. In contrast, the long closed time increased as a function of decreased Ca\(^{2+}\) concentration. In addition, the model correctly predicted the changes in the relative amplitudes of open- and closed-time components.

The low open probability behavior was modeled separately using the same gating scheme (Fig. 11). The best agreement with the experimental data was obtained if all Ca\(^{2+}\)-dependent rates were reduced by a factor of 6.7. The model for low open probability behavior predicted dwell times similar to experimentally derived values for the two Ca\(^{2+}\) concentrations used in the simulation (see Figs. 7 and 11). It also reflected the

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Figure 9. Calcium dependence of kinetic parameters describing low open probability behavior. Data are from three single channel patches except in 0.2 \(\mu\)M Ca\(^{2+}\) (one patch) and in 0.6 \(\mu\)M Ca\(^{2+}\) (two patches) during periods of low open probability activity. (A) Average short and long open times were independent of intracellular Ca\(^{2+}\) concentration. Linear regression analysis of the data is shown as the solid lines and illustrates the Ca\(^{2+}\) independence of the open-time constants. (B) Average short and intermediate closed times were independent of Ca\(^{2+}\) concentration. Solid lines represent linear regression analysis. (C) The long closed times during low open probability activity was Ca\(^{2+}\) dependent, similar to what was observed during high channel open probability (compare Fig. 5). Linear regression analysis yielded a slope of \(-2,460 \text{ ms } \mu\text{M}^{-1}\). (D) The relative amplitude of the short open-time component decreased as a function of Ca\(^{2+}\) concentration. Data are shown on a semi-logarithmic scale for comparison with Fig. 5 (high open probability behavior). The line in (D) is included for display purposes only. (E) The relative amplitude of the short closed-time component increased with increasing Ca\(^{2+}\) concentration. (F) Increasing Ca\(^{2+}\) concentrations lead to a decrease in the relative amplitude of the long closed-time component.
changes seen with changing Ca\(^{2+}\) concentration. A possible shortcoming of the modified model was that it predicted a somewhat larger fraction of intermediate compared with short closures than was observed experimentally. However, this discrepancy was most prominent in low Ca\(^{2+}\), where openings were rare with the obtained histograms containing a relatively small number of events.

The presented gating schemes predict that both long- and short-duration openings should be grouped; i.e., after a long opening, the channel is more likely to reenter the same long open state rather than transition into the short open state. The same should be true for short openings. Consistent with this prediction, long openings were predominantly followed by other long openings in all three patches examined. Fig. 12 A shows histograms containing all openings (left), openings after openings shorter than 1 ms (middle), and openings after openings longer than 10 ms (right) for the patch illustrated in Fig. 4 exposed to 0.4 µM Ca\(^{2+}\). All three histograms were best fit with two exponentials, and the lifetimes of each exponential component were similar under the three conditions. In contrast, the relative amplitudes of the two components varied considerably with the percentage of long openings decreasing from 49% (of all openings) to 20%, when only
openings after short openings were included, and increasing to 74% after long openings. Similar correlations between successive openings were seen in simulations of the model. For the example shown in the insets, 49% of all openings, 23% of openings after short openings, and 81% of openings after long openings were of the long time constant. As illustrated in Fig. 12 B, similar results were obtained for the channel displaying low open probability behavior shown in Fig. 7 (1 μM Ca\(^{2+}\)) and the gating scheme in Fig. 11.

To assure that the low open probability behavior was not a phenomenon limited to oocytes, stably transfected Chinese hamster ovary cells expressing rSK2 channels were examined. Inside-out patches bathed in 0.8 μM Ca\(^{2+}\) contained SK channels that occasionally switched between periods of high and low open probability with open- and closed-time constants not significantly different from the values reported here for patches excised from oocytes (not shown). Therefore, it seemed likely that the low open probability behavior and the switches between different types of channel activity were inherent properties of SK channels rather than a function of the expression system.

The mechanism underlying switches between high and low open probability behavior is unknown, except that the process has some dependence on Ca\(^{2+}\). Since channels reversibly switched behaviors in the absence of added ATP or Mg\(^{2+}\), it seemed unlikely that channel...

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**Figure 12.** Correlations between successive openings. (A) Open-duration histograms for the patch in Fig. 4 exposed to 0.4 μM Ca\(^{2+}\) including all openings (left), openings after openings shorter than 1 ms (middle), and openings after openings longer than 10 ms (right). The lifetimes and relative amplitudes of the two components were 1.1 ms, 51%, and 13.5 ms for all openings; 1.1 ms, 80%, and 15.8 ms after short openings; 1.3 ms, 26%, and 14.0 ms after long openings. Insets show corresponding histograms for simulations of the model in Fig. 10. The short and long components had lifetimes of 1.0 ms, 51%, and 12.2 ms for all openings; 1.0 ms, 77%, and 12.3 ms after short openings; and 1.1 ms, 19%, and 12.8 ms after long openings. (B) Open-duration histograms for the channel in Fig. 7 displaying low open probability behavior in 1.0 μM Ca\(^{2+}\). Open times were fit to two components with lifetimes of 0.9 ms, 71%, and 9.8 ms (left). Openings after openings shorter than 1 ms (middle) had lifetimes of 0.7 ms, 92%, and 8.8 ms. Openings after openings longer than 10 ms (right) were predominantly long but could not be fit due to the small number of events. Insets show corresponding histograms for the model in Fig. 11. The two components had lifetimes of 1.0 ms, 73%, and 11.4 ms for all openings; 1.0 ms, 91%, and 9.1 ms after short openings; and 1.2 ms, 24%, and 13.3 ms after long openings.
phosphorylation plays a role, and the two behaviors may reflect intrinsic conformational states of the channel. The Ca dependence of the time channels spent exhibiting each behavior (Fig. 7 A) suggested that switches between the two behaviors could contribute to the Ca\(^{2+}\) dependence of the macropatch current. This was illustrated by the calculation of an effective open probability in Fig. 7 B, which was right-shifted and somewhat steeper than the single channel open probability.

Comparison with Gating Schemes and Mode Shifts Seen with Other Channels

The gating scheme presented here is the first attempt at modeling the Ca\(^{2+}\)-dependent activation of SK channels. The model shares some similarities with gating schemes used to describe the voltage- and Ca\(^{2+}\)-dependent properties of BK channels in the major gating mode. In particular, the model proposed by DiChiara and Reinhart (1995) for cloned drosophila and human BK channels can be thought of as an expanded version of this model containing an additional closed state connected to a third open state. In addition, DiChiara and Reinhart (1995) also found that only the (three) longest closed states were Ca\(^{2+}\) dependent. However, studies on BK channels from rat skeletal muscle found two of three open times to be Ca\(^{2+}\) dependent and proposed a model in which open states were connected by Ca\(^{2+}\)-dependent transitions (McManus and Magleby, 1991). Earlier studies on rat muscle BK channels incorporated into lipid bilayers (Moczydlowski and Latorre, 1983) and cultured rat muscle cells (Magleby and Pallotta, 1983) also found a Ca\(^{2+}\) dependency of the mean open time. A simple sequential model containing two closed and two open states was proposed in the former and a branched model containing three closed and two parallel open states in the latter study. Differences between these models may at least partially be due to differences in the time resolution. The time resolution used in this study was the same as that used by Magleby and Pallotta (1983).

In addition to the normal mode of gating, four kinetic modes have been observed for BK channels in rat skeletal muscle (McManus and Magleby, 1988; Rothberg et al., 1996). All of them are characterized by a lower channel open probability compared with the normal mode, but together only account for <10% of channel openings at Ca\(^{2+}\) concentrations below 10 \(\mu\)M. Three of the lower open probability modes, termed the intermediate open, brief open, and buzz mode, may result from excluded entry into certain states of the scheme used to describe normal mode gating (McManus and Magleby, 1988). In contrast, the SK channels in this study entered a low open probability mode that appeared to have the same number of open and closed states as the high open probability mode. The frequency and duration of sojourns to the low activity mode of BK channels increased with increasing Ca\(^{2+}\) concentration (Rothberg et al., 1996), while the low open probability mode of recombinant SK channels was entered more frequently in low Ca\(^{2+}\) concentrations.

In contrast to the mode shifting in native BK channels, which occurred rapidly and was followed by periods of stable activity (McManus and Magleby, 1988), cloned *Drosophila* BK channels expressed in oocytes displayed large slow fluctuations between changing levels of open probability (Silberberg et al., 1996). This behavior was termed wanderlust kinetics. Except for the very slow time course of the fluctuations, wanderlust kinetics appears similar to the changes in open probability reported here for cloned SK channels. During low activity periods, recombinant BK and SK channels show bursts of activity with kinetics similar to those seen during high activity. These bursts are separated by long closures not seen during periods of high activity. For both channels, the main difference between time constants describing high and low activity behavior was in the longest closed-time component(s). Silberberg et al. (1996) suggested that wanderlust kinetics can be described by a single gating scheme if one assumes one or more multiplicative factors modifying the Ca\(^{2+}\)-dependent rate constants.

Low activity modes have been observed for a number of other ion channels. These include glutamate-activated channels from locust muscle (Patlak et al., 1979), *Aplysia* voltage-gated cation channels (Wilson and Kaczmarek, 1993), cardiac calcium channels (Hess et al., 1984), frog skeletal muscle sodium channels (Patlak et al., 1986), acetylcholine receptors in frog muscle (Auerbach and Lingle, 1986), M-channels in bullfrog sympathetic neurons (Marriot, 1993), and N-methyl-D-aspartate receptors (Donnelly and Pallotta, 1995). The gating modes observed for the *Aplysia* voltage-gated cation channel appeared most similar to those of cloned SK channels in that the two modes differ mainly by the presence or absence of long closures, whereas the open and the shorter closed times are similar for the two modes.

SK Channel Kinetics May Account for the Time Course of Apamin-sensitive AHPs

The gating scheme and rate constants presented in Fig. 10 could be used to predict the activation time constant of a macroscopic SK current elicited by a fast jump in Ca\(^{2+}\) concentration. A current trace evoked by a jump from 0 to 9.5 \(\mu\)M Ca\(^{2+}\) is shown in Fig. 13 A, superimposed with a simulation of the gating scheme in Fig. 10. The current activated with a time constant of 4.9 ms and, upon removal of free Ca\(^{2+}\), deactivated with a time constant of 58 ms. Averages from six Ca\(^{2+}\) jumps (two patches) were 4.1 ± 0.6 ms for the activation and 57.3 ±
11.1 ms for the deactivation time constant. Simulations of the model (Fig. 10) are shown in Fig. 13 B for jumps to 0.2, 0.5, 1, 5, and 10 μM Ca²⁺. Since macroscopic currents would be dominated by channels displaying high open probability behavior, only this type of channel behavior was considered here. Fitting of the simulated traces to single exponentials yielded activation time constants of ~5 ms in 5 μM and 50 ms in 0.5 μM free Ca²⁺. Figure 13 C shows a plot of the activation rate (τ⁻¹) versus Ca²⁺ concentration. Over the range of 0.2 to 10 μM Ca²⁺, the model predicted an approximately linear relationship between the activation rate of the macroscopic current and the Ca²⁺ concentration with a forward rate constant of 47 μM⁻¹ s⁻¹. This rate was somewhat slower than the slowest Ca²⁺-dependent rate (80 μM⁻¹ s⁻¹) of the gating scheme in Fig. 10, possibly reflecting the requirement for several Ca²⁺-dependent transitions before channel opening. An apparent Ca²⁺-binding rate of 26 μM⁻¹ s⁻¹ was estimated from the macroscopic current activation (Fig. 13 C, ○). A rate of 21 μM⁻¹ s⁻¹ was seen in rat sympathetic neurons using flash photolysis of photosensitive Ca²⁺ chelators (Gurney et al., 1987). Both rates are fast and, considering the uncertainty associated with access of Ca²⁺ to the patch (this study) or with determinations of the amount of released Ca²⁺ (Gurney et al., 1987), are in reasonable agreement with our model. In the study of Gurney et al. (1987), a lower limit of 25 s⁻¹ for channel closure upon Ca²⁺ removal was estimated by extrapolation of the data to 0 Ca²⁺. This rate was somewhat faster than, but not markedly different from, the 17 s⁻¹ measured during rapid Ca²⁺ removal (τ = 58 ms, Fig. 13 A) and the 13 s⁻¹ (τ = 76 ms, Fig. 13 B) predicted by our model. In comparison, the apamin-sensitive component of the AHP in bullfrog sympathetic ganglion cells decayed with a time constant of 150 ms, corresponding to a rate of 7 s⁻¹ (Lancaster and Pennefather, 1987).

SK channels underlie the AHP in many neurons. The apamin-sensitive component of the AHP in vagal motorneurons (Sah, 1992) and bullfrog sympathetic ganglion (Pennefather et al., 1985) activates rapidly and is maximal <5 ms after the voltage step used to evoke an action potential. This time course suggests that SK channels in these cells are located near voltage-dependent calcium channels. In contrast, the apamin-insensitive component of the AHP has a slower time course activating with a time constant of 550 ms in guinea pig vagal neurons (Sah, 1993). The apamin-insensitive member (SK1) of the cloned SK channel subtypes is highly homologous to apamin-sensitive members (SK2 and SK3) and shares the same Ca²⁺ sensitivity (Köhler et al., 1996). As expected, preliminary experiments show that hSK1 and rSK2 have similar open and closed times. Stationary and nonstationary fluctuation analysis of the current at the peak of the apamin-insensitive slow AHP yielded open probabilities of 0.4 in hippocampal pyramidal neurons (Sah and Isaacson, 1995), 0.6 in dentate granule neurons of the hippocampus (Valiante et al., 1997), and 0.7 in vagal motoneurons (Sah, 1995). Assuming that apamin-sensitive and insensitive SK channels possess similar single channel kinetics, this suggests that the effective Ca²⁺ concentration at the channel was in the range of 0.7–1.2 μM. In this concentration of Ca²⁺, the model for the high open probability behavior were well described by single exponentials, yielding time constants of 4.9 and 58 ms. Overlaid on the data is a simulation of the model in Fig. 10. (B) Computer simulations of the gating scheme in Fig. 10 for 400 ms Ca²⁺-jumps from 0 to the Ca²⁺ concentration indicated above each trace. For each trace, the rising and falling phase of the activation and deactivation time course, respectively, were fit to single exponential functions with the fitted lines coinciding with the traces. The deactivation time course was independent of the Ca²⁺ concentration during the jump and gave a time constant of 76 ms. (C) Ca²⁺ dependence of the activation time course. Activation time-constants obtained from fitting the macroscopic current activation (○) and the simulated traces in B (●) were plotted as a function of Ca²⁺ concentration. Over the range of 0.2 to 10 μM Ca²⁺, the Ca²⁺ dependence of the simulated data appeared linear and regression analysis yielded an apparent Ca²⁺-binding rate of 47 μM⁻¹ s⁻¹.
predicts an activation time constant of 20–30 ms, >10-fold faster than the time constant of the apamin-insensitive slow AHP in vagal neurons (Sah, 1993). This suggests that SK channel kinetics alone are insufficient to explain the time course of the apamin-insensitive AHP, in agreement with data from hippocampal pyramidal neurons using flash photolysis of caged Ca$^{2+}$ (Lancaster and Zucker, 1994).

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