Consequences of the Stoichiometry of Slo1 α and Auxiliary β Subunits on Functional Properties of Large-Conductance Ca2+-Activated K+ Channels

Ying-Wei Wang, Jiu Ping Ding, Xiao-Ming Xia, and Christopher J. Lingle

Departments of Anesthesiology and Anatomy and Neurobiology, Washington University School of Medicine, St. Louis, Missouri 63110

Auxiliary β subunits play a major role in defining the functional properties of large-conductance, Ca2+-dependent BK-type K⁺ channels. In particular, both the β1 and β2 subunits produce strong shifts in the voltage dependence of channel activation at a given Ca2+. β subunits are thought to coassemble with α subunits in a 1:1 stoichiometry, such that a full ion channel complex may contain up to four β subunits per channel. However, previous results raise the possibility that ion channels with less than a full complement of β subunits may also occur. The functional consequence of channels with differing stoichiometries remains unknown. Here, using expression of α and β subunits in Xenopus oocytes, we show explicitly that functional BK channels can arise with less than four β subunits. Furthermore, the results show that, for both the β1 and β2 subunits, each individual β subunit produces an essentially identical, incremental effect on the voltage dependence of gating. For channels arising from α + β2 subunits, the number of β2 subunits per channel also has a substantial impact on properties of steady-state inactivation and recovery from inactivation. Thus, the stoichiometry of α:β subunit assembly can play a major functional role in defining the apparent Ca2+ dependence of activation of BK channels and in influencing the availability of BK channels for activation.

Key words: auxiliary subunits; BK channels; Ca²⁺- and voltage-gated K⁺ channels; Slo1 channels; inactivation; ion channel stoichiometry; gating mechanisms

Large-conductance, Ca²⁺-activated BK-type K⁺ channels exhibit substantial functional diversity (McManus, 1991; Vergara et al., 1998) contributed, in part, from coexpression of the pore-forming Slo α subunit (Adelman et al., 1992; Butler et al., 1993) with members of an auxiliary β subunit family. At present, four mammalian β subunits have been identified (Knaus et al., 1994b; Wallner et al., 1999; Xia et al., 1999, 2000; Brenner et al., 2000; Meera et al., 2000; Uebele et al., 2000; Weiger et al., 2000). Both the β1 and β2 subunits result in pronounced negative shifts in the voltage of half-activation at a given [Ca²⁺] (McManus et al., 1995; Wallner et al., 1995, 1999; Xia et al., 1999; Brenner et al., 2000). Both the β2 (Wallner et al., 1999; Xia et al., 2000) and β3b (Uebele et al., 2000; Xia et al., 2000) subunits result in kinetically distinct inactivating BK channels.

β subunits can exist in a 1:1 stoichiometry with α subunits (Knaus et al., 1994a): four β subunits can coassemble with four α subunits into an intact BK channel. Previous work on inactivating BK (BKc) channels in rat chromaffin cells (Ding et al., 1998) suggests that the variability in inactivation behavior might arise from differential stoichiometry of some inactivation-competent subunit in the channel population (Ding et al., 1998). Given the presence of β2 subunit message in rat chromaffin cells (Xia et al., 1999) and the similarity of α + β2 currents to BKc currents (Wallner et al., 1999; Xia et al., 1999), one possibility is that, in rat chromaffin cells, channels occur with less than a 1:1 assembly of β2:α subunits. The dependence of BK channel properties on β:α coassembly was also examined in Xenopus oocytes by varying the ratio of coinjected β1 and α subunits (Jones et al., 1999). This work proposed the view that β1 subunits produced an all-or-none shift in gating properties of the resulting BK channels (Jones et al., 1999).

These previous studies raise interesting questions concerning the functional consequences that result from less than a full 1:1 stoichiometric assembly of β and α subunits BK. First, it remains unclear whether BK channels can form with less than four β subunits. Second, if BK channels can contain less than four β subunits, what is the role that a single β subunit plays in influencing the various functional properties of the channel? To address these issues, we use the inactivation properties conferred on BK channels by the β2 subunit as an indicator of β2:α subunit stoichiometry within a channel population that can then be related to other functional properties. The results demonstrate that BK channels that contain less than four β subunits can occur.

Furthermore, channels with less than a full complement of β subunits show gating properties and inactivation behavior that scale with the average number of β subunits per channel.

MATERIALS AND METHODS

Expression in Xenopus oocytes. The preparation of the β1 and β2 expression constructs used here has been described previously (Xia et al., 1999). Two other constructs used here were also described in previous work (Xia et al., 1999): first, the β2-Δ33 construct in which 33 N-terminal amino acids were removed from the β2 subunit; and second, a construct in which the 33 initial amino acids from the β2 N terminus were appended to the N terminus of the β1 subunit. This latter construct is here termed β1–C2. The α subunit used here was the mouse Slo1 construct used previously (Xia et al., 1999), which corresponds to a zero amino acid insert at splice site 1 and a three amino acid insert at splice

Received Nov. 2, 2001; revised Dec. 7, 2001; accepted Dec. 7, 2001.

This work was supported by National Institutes of Health Grants DK46564 and NS37671. We thank Lynn Lavack for injection and care of oocytes.

Correspondence should be addressed to Chris Lingle, Department of Anesthesiology, Washington University School of Medicine, Box 8054, St. Louis, MO 63110.

E-mail: clingle@morpheus.wustl.edu.

Copyright © 2002 Society for Neuroscience 0270-6474/02/221550-12$15.00/0

Key words: auxiliary subunits; BK channels; Ca²⁺- and voltage-gated K⁺ channels; Slo1 channels; inactivation; ion channel stoichiometry; gating mechanisms

Expression in Xenopus oocytes. The preparation of the β1 and β2 expression constructs used here has been described previously (Xia et al., 1999). Two other constructs used here were also described in previous work (Xia et al., 1999): first, the β2-Δ33 construct in which 33 N-terminal amino acids were removed from the β2 subunit; and second, a construct in which the 33 initial amino acids from the β2 N terminus were appended to the N terminus of the β1 subunit. This latter construct is here termed β1–C2. The α subunit used here was the mouse Slo1 construct used previously (Xia et al., 1999), which corresponds to a zero amino acid insert at splice site 1 and a three amino acid insert at splice
yielded values essentially identical to our own Cl/H11002 (in mM): 140 potassium methanesulfonate, 20 KOH, 10 HEPES, pH 7.0, solutions bathing the cytoplasmic face of the patch membrane contained channels were assumed to contain four possible units follows that outlined in previous work (Ding et al., 1998). All Slo1 species. The molecular weight of Slo1 in Xenopus oocytes, even when we are reasonably confident that RNA degradation has been minimized. To minimize degradation problems that might be associated with freezing and thawing, each preparation of RNA was separated into aliquots at the time of preparation, and a separate aliquot was used for each injection. Another potential problem is that, for distinct nonhomologous RNA species (i.e., α and β eRNA), it is not clear how the relative abundance may relate to the stoichiometry of assembly. To circumvent this problem, we have therefore used the properties of inactivation as independent estimators of the stoichiometry of channel assembly.

Electrophysiology. Macroscopic and single-channel current measurement follow methods in standard use in this laboratory. For these experiments, currents were recorded in the inside-out patch configuration (Hamill et al., 1981). Digitization for macroscopic currents was typically at 10–50 kHz with analog filtering during acquisition (5–20 kHz, Bessel low-pass filter, −3 dB). For single-channel experiments, digitization was at 100 kHz, with 5 kHz filtering. Preparation of the pipette solution and Ca2+ solutions has been described previously (Wei et al., 1994; Xia et al., 1999). The pipette extracellular solution was (in mM): 140 potassium methanesulfonate, 20 KOH, 10 HEPES, and 2 MgCl2, pH 7.0. Test solutions bathing the cytoplasmic face of the patch membrane contained (in mM): 140 potassium methanesulfonate, 20 KOH, 10 HEPES, pH 7.0, and one of the following: 5 mM EGTA (for nominally zero Ca2+ and 0.5 and 1 μM Ca2+ solutions), 5 mM HEDTA (for 4 and 10 μM Ca2+ solutions), or no added Ca2+ buffer (for 60, 100, and 300 mM CaCl2 and 1 and 5 mM Ca2+ solutions). The methanesulfonate solutions were calibrated against a commercial set of Ca2+ standards (WPI, Sarasota, FL), which yielded values essentially identical to our own Cl/H11002/H9251/H11001/H9251/H11001 based standards. Local perfusion of membrane patches was as described previously (Soloro and Lingle, 1992; Soloro et al., 1997).

pClamp 7.0 or pClamp 8.0 for Windows (Axon Instruments, Foster City, CA) was used to generate voltage commands and to digitize currents. Current values were measured using ClampFit (Axon Instruments), converted to conductances, and then fit with a custom nonlinear least squares fitting program. Conductance–voltage (G–V) curves for activation were fit with a Boltzmann equation with the form:

\[ G(V) = G_{\text{max}} \times \left( 1 + \exp\left(\frac{V - V_{\text{shift}}}{k}\right) \right)^{-1}, \]

where \( G_{\text{max}} \) is the voltage of half-maximal activation of conductance, and \( k \) is the voltage dependence of the activation process (mV−1). Experiments were done at room temperature (21–24°C). All salts and chemicals were obtained from Sigma (St. Louis, MO).

Simulation of G–V curves based on partial occupancy of β2 subunit binding sites. The strategy for evaluation of the functional consequences of channel populations containing differing stoichiometries of β:α subunits follows that outlined in previous work (Ding et al., 1998). All channels were assumed to contain four possible β subunit binding sites. Fractional occupancy by β subunits of those sites was assumed in all cases to follow a binomial distribution. At a given fractional occupancy, the fraction of channels in any of the possible stoichiometries was then calculated, and the contribution of channels of a particular stoichiometry to the overall G–V curves was determined based on different assumptions (e.g., independence, positive cooperativity, or negative cooperativity) about β subunit effects (Ding et al., 1998). Time constants for inactivation of a channel population containing α + β2 subunits in differing stoichiometries were expected to exhibit up to four exponential components (corresponding to the presence of one to four inactivation domains). However, empirically, the relative amplitudes and time constants of these components result in currents that decay with a time course that can be reasonably approximated by a single exponential (Ding et al., 1998). To generate predictions for the inactivation time constant for channel populations containing some average number of β2 subunits per channel, currents were simulated and fit with single exponentials.

RESULTS

Inactivation properties of α + β2 currents indicate that channels can contain less than four inactivation domains and that inactivation domains act in an independent manner

Previous work has suggested that the inactivation properties of native inactivating BK currents among different chromaffin cells can be used as indicators of the average stoichiometry of assembly of inactivating and noninactivating subunits (Ding et al., 1998). At least in regard to inactivation behavior, each inactivating subunit appears to behave in an independent manner. Thus, the average number of inactivating subunits per channel within a population of channels defines the average inactivation rate of channels in that population. With identification of the β2 auxiliary subunit in chromaffin cells (Xia et al., 1999), this raised the possibility that variability in the average number of β2 subunits (or other inactivating β subunit) per channel may account for the previous observations in chromaffin cells.

If, in fact, inactivation properties and \( \tau_i \) provide a direct assay for the stoichiometry of β2 α subunits in a channel population; it therefore becomes possible to examine the consequences of subunit stoichiometry on other channel functional properties without having specific information about the expression levels of subunits within the cell. This is particularly advantageous when it is unclear to what extent oocyte-to-oocyte variability or variability in RNA preparations may have an impact on the ability of subunits to be expressed. Using this strategy, we have therefore sought to address how channel stoichiometry may affect other functional properties of the resulting BK channels.

Specifically, the β2 subunit was co-injected with mSlo α subunits into Xenopus oocytes at different ratios, and the following aspects of BK channel function were determined: (1) the relationship between conductance and activation voltage at 10 and 300 μM Ca2+; (2) the rates of onset and recovery from inactivation; (3) the ratio of inactivating to noninactivating current; and (4) the voltage dependence of steady-state inactivation.

Figure 1 shows families of currents activated by depolarizing voltage steps at either 10 or 300 μM Ca2+ for four different injection ratios of β2 and α subunits. Qualitatively, as the relative amount of β2 subunit is reduced, there is less current activation at potentials negative to zero, \( \tau_i \) is slowed, and there is a larger noninactivating component of current at the end of the most depolarized voltage step. All of these changes are those expected for a model in which various indicators of BK channel function scale in accordance with the average number of β2 subunits per channel. This is examined more explicitly below. Another feature of the currents shown in Figure 1 is that peak current activated at positive command potentials is smaller with 300 than with 10 μM Ca2+. This reflects the persistence of steady-state inactivation even after a 100 msec step to −180 mV.

The slowing of \( \tau_i \) as a function of the injected ratio of β2:α subunits is illustrated in Figure 2A for currents obtained with 300 μM Ca2+ at either +100 or +160 mV. \( \tau_i \) reaches a limit of ~20 msec, at ratios of both 1:1 and 2:1 suggesting that maximal occupation of α subunits by β2 subunits has occurred. The slowest observed values of \( \tau_i \) are ~90 msec. This value is a bit larger than the theoretical limit of 80 msec predicted for an inactivation model involving four independent inactivation domains, in which
Figure 1. Decreasing the ratio of injected β2:α subunits slows the inactivation time constant of α + β2 currents. A–E, Traces show currents obtained in inside-out patches, with each patch from an oocyte injected with the indicated ratio of β2:α subunits. From top to bottom, traces correspond to oocytes injected with 1:1 β2:α (A), 0.05 β2:α (B), 0.025 β2:α (C), 0.01 β2:α (D), and α alone (E). Left traces were obtained in 10 μM Ca²⁺, and right traces were obtained in 300 μM Ca²⁺. Traces show currents activated to potentials between −100 and +120 mV in steps of 20 mV, with tail currents at −120 mV with a prepulse to −180 mV. The reduction in peak current activation with 300 μM Ca²⁺ corresponds to the additional steady-state inactivation of channels at −180 mV.

20 msec is the minimal \( \tau_i \). However, measurement of the slowest \( \tau_i \) values can be influenced by other factors. For example, at positive activation potentials, Slo1 currents, even in the absence of β subunits, can exhibit a slow reduction in current during depolarization (Fig. 1E). The presence of such additional slow blocking components at +100 and +160 mV would tend to slow the apparent inactivation time constant resulting from β2 subunit action, which might account for the slower than expected time constants observed at the 0.01 β2:α injection ratio.

The voltage dependence of \( \tau_i \) at both low (0.025) and high (1.0) ratios of β2:α subunits is plotted in Figure 2B. Because it is known that β subunits shift the voltage dependence of activation at a given [Ca²⁺], a shift in \( \tau_i \) might occur simply because of a shift resulting from coupling of inactivation to activation. However, over the range of +100 to +160 mV, there is little voltage dependence to \( \tau_i \) at either injection ratio. This indicates that the large changes in \( \tau_i \) at +160 mV that are observed as a consequence of different injection ratios must reflect the underlying stoichiometry of the inactivation process and not a consequence of a shift in activation potentials.

As mentioned in Materials and Methods, the injection ratio of β2:α subunits does not provide any handle on the stoichiometry of assembly within the oocyte. Therefore, we have attempted to use the inactivation behavior to reveal something about channel stoichiometry. As above, we measured \( \tau_i \) during activation steps to +160 mV at 10 μM Ca²⁺. For the same currents, we also measured the peak current activated by the voltage step to +160 mV and the steady-state current at the end of the voltage step (300 msec). For the simple model in which channel stoichiometries are
defined by a binomial distribution and up to four β2 subunits independently contribute to the onset of inactivation (MacKinnon et al., 1993; Ding et al., 1998), the ratio of steady-state current to peak current ($f_{ss}$) should vary in accordance with $\tau_i$; at the largest steady-state current, $\tau_i$ should reach a limiting value that is approximately fourfold slower than the fastest values of $\tau_i$. The conditions for these experiments were chosen for the following reasons. At $10 \mu M$ Ca$^{2+}$, there is minimal steady-state inactivation at a potential of $-140$ mV, so that a subsequent activation step should define the maximal current expected for the total population of expressed channels (Ding and Lingle, 2002); +160 mV was used as an activation step so that the kinetics of channel opening at $10 \mu M$ Ca$^{2+}$ are relatively fast compared with the onset of inactivation. A drawback of the use of a step to $+160$ mV is that there are usually slow “inactivation” components, perhaps because of a divalent cation block that may contaminate the estimate of $\tau_i$ and steady-state current. This latter issue is most problematic for the smaller $\beta 2: \alpha$ ratios.

The relationship between changes in $\tau_i$ as a function of $f_{ss}$ is plotted in Figure 2C for several $\beta 2: \alpha$ injection ratios. Over the range of injection ratios used, $\tau_i$ ranged from $\sim 20$ msec to time constants of $>80$ msec. The basic heteromultimeric model for Shaker K$^+$ inactivation (MacKinnon et al., 1993) and BK channel inactivation (Ding et al., 1998) would predict that, at the limit of the lowest ratios of $\beta 2: \alpha$ subunits, $\tau_i$ should approach approximately four times that at the highest ratios. The values exhibit considerable scatter but follow the general trend required by an inactivation model in which up to four $\beta 2$ subunits, each with an independently acting inactivation domain, can contribute to an intact channel containing $\alpha$ + $\beta 2$ sublines. Lines are drawn over the data showing the predictions for this model for cases in which the minimal $\tau_i$ is 17.5–25 msec. At the lowest $f_{ss}$, the results appear to follow the expectations for the lines corresponding to minimal $\tau_i$ values of $\sim 20$ msec. As $f_{ss}$ increases, values for $\tau_i$ deviate from the theoretical expectations. This was also observed in rat chromaffin cells during trypsin-mediated removal of inactivation (Ding et al., 1998). The values at high $f_{ss}$ are likely to be in error for two reasons. The slow blocking processes mentioned above will result in slower values of $\tau_i$, and any additional slow blocking processes will also reduce the value of $f_{ss}$. Both errors will contribute to the observed deviation at the lower injection ratios. However, the change in $\tau_i$ relative to $f_{ss}$ argues strongly that channels can assemble with less than a full complement of $\beta 2$ subunits and that each subunit acts in an independent manner to contribute to the onset of inactivation.

Each $\beta 2$ subunit contributes incrementally to the shift in activation $V_{0.5}$

To examine the dependence of the activation of conductance on various injection ratios, $G$–$V$ curves (for $10 \mu M$, Fig. 3d; for $300 \mu M$, Fig. 3B) were calculated from measurement of peak currents. As the $\beta 2: \alpha$ ratio is reduced, $V_{0.5}$ at either 10 or $300 \mu M$ Ca$^{2+}$ is shifted to more positive potentials approaching the values for $\alpha$ alone at the lowest $\beta 2: \alpha$ ratios. For this set of patches, the full shift in $G$–$V$ curves resulting from the $\beta 2$ subunits was 77.8 mV at $10 \mu M$ and 66.3 mV at $300 \mu M$.

There are two possible explanations for the action of $\beta$ subunits that might affect the shift in $G$–$V$ curves. In one case, as the mole fraction of $\beta 2$ subunits is reduced, the fraction of channels containing less than four $\beta 2$ subunits will be increased. The shift in $V_{0.5}$ could arise from channels with less than a full complement of $\beta 2$ subunits having a smaller shift in $V_{0.5}$ than those with four $\beta 2$ subunits. In this case, the $G$–$V$ curves would represent a binomially weighted sum of five distinct Boltzmann functions corresponding to the five possible $\beta 2: \alpha$ stoichiometries (Fig. 3C). Alternatively, it is possible that the shift in the $G$–$V$ curves arises from changes in the proportion of two functional populations of channels, each with a characteristic $V_{0.5}$. Channels containing one or more $\beta 2$ subunits would all share a similar $V_{0.5}$, whereas those containing no $\beta 2$ subunits would appear as $\alpha$ alone. If so, the $G$–$V$ curves would represent a weighted sum of two Boltzmann functions (Fig. 3D). This is the mechanism implied by observations in one previous study (Jones et al., 1999). This latter model might also appear to arise when there is strong cooperativity in the channel assembly process, such that channels contain either four or zero $\beta 2$ subunits.

Comparison of the expectations arising from each type of model with the actual $G$–$V$ curves obtained at $10 \mu M$ Ca$^{2+}$ (Fig. 3A) suggests that a model in which each $\beta 2$ subunit exerts some incremental contribution to the processes involved in shifting the $G$–$V$ curve better approximates the actual results. We also considered two other cases: first, a case of strong positive cooperativity in which each additional $\beta 2$ subunit associated with a channel results in a stronger effect on the $V_{0.5}$ (Fig. 3E); and second, a case of strong negative cooperativity in which most of the shift in $V_{0.5}$ results from the action of a single $\beta 2$ subunit, with smaller effects contributed by each additional $\beta 2$ subunit (Fig. 3F). The latter case was essentially indistinguishable from the case in which a single $\beta 2$ subunit accounted for all the shift in $V_{0.5}$.

The $V_{0.5}$ for activation of conductance shifts in accordance with the fraction of injected $\beta 2$ subunit (Fig. 4/4). Similar to the relationship between $\tau_i$ and $\beta 2: \alpha$ ratio, the $V_{0.5}$ reaches a limiting value at ratios of 1.0 and 2.0. Values for $V_{0.5}$ at a ratio of 0.01 approach those for Slo1 $\alpha$ alone. Because there is no simple relationship between the injected ratio of $\beta 2: \alpha$ subunits and the resulting channel stoichiometry, we have used the relationship between $\tau_i$ and $V_{0.5}$ to examine the effect of stoichiometry on activation $V_{0.5}$. In Figure 4B, the activation $V_{0.5}$ measured at $10 \mu M$ Ca$^{2+}$ is plotted as a function of $\tau_i$. Making a specific assumption about the minimal $\tau_i$, $\tau_i$ can then be used to make estimates of the average number of $\beta 2$ subunits per channel. Vertical lines correspond to the expected time constants for a channel population with an average of four, three, two, and one $\beta 2$ subunits per channel. It should be realized that, except in the case of four, these expected values for the time constants are not equivalent to those predicted when all channels contain a given number of $\beta 2$ subunits. This analysis suggests that the values of $\tau_i$ obtained in these experiments probably reflect average stoichiometries that vary from approximately four $\beta 2$ subunits per channel to less than one $\beta 2$ subunit per channel. Figure 4B also shows the predicted relationship between $V_{0.5}$ and $\tau_i$ expected when each $\beta 2$ subunit contributes an identical amount of shift in $V_{0.5}$. The two lines compare predictions for the cases in which $\tau_i$ is 20 and 25 msec.

Given the scatter in experimental estimates and the variation in $V_{0.5}$ that seems to naturally occur among different sets of experiments, caution must be taken in attempting to relate how much shift in $V_{0.5}$ may occur in accordance with which levels of occupancy of the channels by $\beta 2$ subunits. However, on the basis of the model of inactivation in which the average $\tau_i$ in a patch reflects some average, binomially distributed occupancy of channels by $\beta 2$ subunits, we can calculate a predicted average number of $\beta 2$ subunits per channel, relating it to the observed values for activation $V_{0.5}$ (Fig. 4C). The estimate of $\beta 2$ subunits per channel depends on the assumption of a particular value for $\tau_i$ and the
predicted relationship for $\tau_{\text{min}}$ values of either 20 or 25 msec is illustrated. This procedure suggests that $V_{0.5}$ appears to shift in an approximately linear manner with the number of $\beta_2$ subunits per channel between the limits defined by $\alpha$ subunits alone and full occupancy by $\beta_2$ subunits. For comparison, predictions based on models with either positive or negative cooperativity (from Fig. 3E,F) are also shown in Figure 4C, indicating that the actual results fit better with the model in which each subunit adds linearly to shift the gating equilibrium of the resulting $\alpha + \beta_2$ channels.

In sum, these results are most consistent with the view that each $\beta_2$ subunit independently contributes a fixed amount to the shift in activation $V_{0.5}$.

**Single $\alpha + \beta_2$ channels exist in four different channel stoichiometries, each with a different voltage dependence of activation at a given Ca$^{2+}$**

It might be argued that the failure to observe an all-or-none effect of a single $\beta_2$ subunit reflects the possibility that in macropatch recordings the $G-V$ curves from two functional types of channel (Fig. 3D) simply average to be indistinguishable from the curves predicted for five separate functional types (Fig. 3C). To address this possibility, we therefore turned to single-channel recordings. In Figure 5, example sweeps and ensemble averages from single-channel patches are illustrated, corresponding to channels that inactivated with $\tau_{\text{g}}$ of 21.8, 33.4, 56.4, and 99 msec. A total of 49 single-channel patches were examined. A frequency histogram of the number of occurrences of single-channel inactivation time constants of various values is plotted in Figure 6A. Although the observed values show considerable variability, a fit of a four-component Gaussian distribution indicates that the values cluster at peaks corresponding to 22.6, 33.0, 49.1, and 99.8 msec. Although the number of examples of more slowly inactivating channels is a bit limited, these values correspond quite well to those that would be predicted for four stoichiometries with fully independent inactivation by each inactivation domain (e.g., 22, 33, 41, 51).
spond to cases in which the minimal assumption that the minimum E cooperativity (Fig. 3). Vertical lines 4, 3, 2, and 1 correspond to τ expected for a binomially distributed stoichiometry with an average of four, three, two, and one β2 subunits per channel, respectively, with the minimum τ assumed to be 20 msec. Horizontal lines correspond to V0.5 ± SD recorded for currents resulting from α alone. Dashed lines correspond to an empirical relationship between V0.5 and the average number of β2 subunits per channel calculated from the inactivation time constants. Large ○, Data values from B (obtained at 10 μM Ca2+) with the assumption that the minimum τi for a channel with four β2 subunits is 20 msec. ◆, Same calculation but with a minimum τi of 25 msec. For this conversion, when τi for a patch exceeded the minimal or maximal τi, the average number of β2 subunits per channel was assumed to reflect either four or zero β2 subunits, respectively. For comparison, the V0.5 at different average numbers of β2 subunits are shown for the cases of positive cooperativity (Fig. 3E; ○) and negative cooperativity (Fig. 3F; ◆) and for an additive, incremental effect of each β2 subunit (Fig. 3C; ◆).

44, and 88 msec). These results provide direct evidence that individual channels can contain zero to four β2 subunits per channel.

An interesting aspect of the single-channel recordings was that single-channel current amplitude at +100 mV appeared to scale with the number of β2 subunits per channel, with smaller current amplitudes at higher numbers of β2 subunits (Fig. 5). To confirm that observation, single-channel current measurements were made at potentials from +20 to +100 mV. Single-channel conductances were 214 ± 28.1 pS (τi = 21.6 ± 1.42; three patches), 228 ± 22 pS (τi = 31.3 ± 2.4 msec; three patches), 234 ± 14 pS (τi = 48.2 ± 2.5 msec; three patches), and 278 ± 15 pS (τi = 98.8 ± 1.2 msec; two patches), with extrapolated zero current potentials of <2 mV. The estimate under these conditions for the conductance of α alone is ~270 pS. Thus, the differences in single-channel current amplitudes seen here reflect a true difference in single-channel conductance resulting from the presence of the β2 subunit.

We next addressed the issue of how β2:α subunit stoichiometry affects activation when observed in single channels. To do this, after determination of single-channel τi, inactivation was then removed by brief application of trypsin (0.5 mg/ml) to the cytosolic face of patches. Ensemble averages were then generated at voltages from +20 to +140 mV using 4 μM Ca2+. Four micromolar Ca2+ was preferable to 10 μM Ca2+ for this...
experiment, because with 4 μM Ca^{2+}, the voltage of half-activation is sufficiently positive to 0 mV both for purely α + β2 channels and for α-alone channels to allow better estimation of activation V_{0.5}. Figure 6B shows the resulting estimates of normalized open probability P(0) for such trypsin-treated single-channel recordings along with estimates from four patches containing α subunits alone. The P(0)–V curves span a range of ~60 mV, somewhat similar to that seen with macroscopic currents in Figure 3A. The fitted estimate of V_{0.5} for each single-channel recording is plotted as a function of τ_i in Figure 6C. For this set of 15 single-channel patches, the values appear to cluster into four groups, with the inactivation time constant strongly correlated with the activation V_{0.5}. Coupled with other results presented above, this strongly argues that each β2 subunit independently produces an equivalent effect on the resulting V_{0.5} at a given [Ca^{2+}].

**Figure 6.** Channel stoichiometry revealed by inactivation correlates with an incremental shift in activation V_{0.5}. A, A frequency distribution of τ_i values determined from ensemble current averages for 49 single-channel patches is plotted. τ_i values were distributed into 5 msec bins, and a four-component Gaussian function was fit to the binned data, resulting in peak values of 22.6, 33.0, 49.1, and 99.8 msec, as indicated. For an inactivation mechanism with four independently acting inactivation domains, if channels with only one domain inactivate with a time constant of 100 msec, channels with two to four inactivation domains are predicted to inactivate with τ_i of 50, 33, and 25 msec, respectively. B, A set of the channels studied in A were Briefly treated with trypsin to remove inactivation and the relationship between P(0) and activation potential was determined at 4 μM Ca^{2+}. Each filled symbol corresponds to a different patch expressing a channel with some mixture of α + β2 subunits. Open symbols correspond to patches expressing α alone. C, The relationship between activation V_{0.5} and τ_i is plotted for 15 patches studied as in A and B. Channels that exhibited a slower τ_i exhibited a more positive activation V_{0.5}. Values appear to cluster into four groups, which were chosen by eye and indicated by the different symbols. D, Values in C were grouped as indicated, and V_{0.5} was plotted as a function of 100/τ_i. 100/τ_i should reflect the number of β2 subunits per channel assuming that a channel with one β2 subunit inactivates with τ_i = 100 msec. The value at 0 corresponds to V_{0.5} values for single-channel patches containing α subunits only.

**Figure 7.** Steady-state inactivation shifts with changes in the β2:α subunit ratio. A1–A5, Currents were activated with the voltage protocol indicated on the top for five different β2:α injection ratios with 10 μM Ca^{2+}. At smaller ratios, the amount of noninactivating current increases, and the rate of inactivation slows. The duration of the conditioning step was 600 msec. The dashed lines indicate the 0-current level. B, The percent availability ± SD of only the inactivating portion of current is plotted as a function of conditioning potential for four to six patches at each of the indicated β2:α injection ratios. For ratios of 0.01, 0.025, 0.05, 0.1, 1.0, and 2.0, V_{0.5} values were ~17.2, ~38.9, ~60.1, ~73.1, ~88.5, and ~97.6 mV, respectively. C, The maximal current activated from different conditioning potentials was determined, providing an indication of the percent availability of the total channel population at different potentials. For ratios from 0.01 to 2.0, the voltages at which half the channels in the total population were available for activation were ~97.6, ~88.5, ~67.8, ~29.7, ~3.5, and ~37 mV. D, The voltage of half-availability determined in B is plotted as a function of injection ratio.

**Steady-state inactivation is also dependent on the average number of β2 subunits per channel**

Other physiologically important properties of α + β2 currents may also be dependent on the stoichiometry of channel composition. Therefore, the dependence of two other properties of inactivating BK channels on the ratios of β2:α subunits was also determined: first, the voltage dependence of steady-state inactivation measured at 10 μM Ca^{2+}; and second, the rate of recovery from inactivation at ~140 mV also with 10 μM Ca^{2+}.

To examine steady-state inactivation properties, patches were held for at least 500 msec at conditioning potentials between ~190 and +10 mV (Fig. 7A) before steps to +160 mV. At higher mole fractions of the β2 subunit, steady-state inactivation mea-
Inactivated with 10 μM cytosolic Ca$^{2+}$ was essentially identical to that previously observed for inactivating BK channels in RIN cells (Li et al., 1999), and for BK$\alpha$ channels in chromaffin cells (Ding and Lingle, 2002). The fractional availability of current as a function of the initial conditioning potential was determined. As the ratio of $\beta_2\alpha$ was reduced, the fractional availability of the resulting BK current was shifted to more positive potentials, and at the lowest dilutions it is clear that there is a substantial amount of current that does not inactivate. Because the activation step was to +160 mV, any channel that contains even one inactivation domain will contribute little to the steady-state current. Thus, the steady-state current represents almost exclusively the fraction of channels that contain no $\beta_2$ subunits. The currents were analyzed in two ways. First, we measured only the fractional availability of the inactivating portion of the current (Fig. 7B). This takes into account only those channels that contain at least one $\beta_2$ subunit. We also plotted the total amount of current available from any conditioning potential (Fig. 7C). This provides a better indication of the entire channel population at each $\beta_2\alpha$ dilution. Qualitatively, it is quite clear that reducing the ratio of injected $\beta_2\alpha$ subunits results in two effects: first, a marked shift in the fractional availability of channels for activation; and second, an increase in the fraction of channels that do not inactivate. The dependence of the voltage of half-channel availability on injection ratio is shown in Figure 7D.

**Recovery from inactivation exhibits an anomalous dependence on $\beta_2\alpha$ ratio**

The time constant of recovery from inactivation ($\tau_r$) was defined at 10 μm with a paired pulse protocol in which, after complete inactivation of the channels at +140 mV, a second test step to +140 mV followed a variable recovery interval at −140 mV (Fig. 8A). At high $\beta_2\alpha$ ratios, $\tau_r$ was 20–25 msec similar to values measured for inactivating BK channels in RIN cells (Ding et al., 1998; Li et al., 1999). As the ratio of $\beta_2\alpha$ was reduced (Fig. 8B, C), $\tau_r$ became faster, appearing to reach a limiting value at ~5 msec at the smallest ratios of $\beta_2\alpha$.

This change of $\tau_r$ with the ratio of $\beta_2\alpha$ would appear to contradict previous work in which progressive trypsin-mediated removal of inactivation of BK channels in chromaffin cells did not alter $\tau_r$ (Ding et al., 1998). For a model of a block in which occupancy of a blocking site by a single inactivation domain is sufficient to produce inactivation, if recovery is governed solely by dissociation of a single domain from its blocking site, no alteration in $\tau_r$ is expected as the number of inactivation domains per channel is altered. Thus, the present result would seem to conflict with previous results and may challenge one simple conception of the molecular steps involved in the inactivation process, namely that the recovery process should be governed by dissociation by a single inactivation domain from its blocking site.

Is there a possible explanation for the dependence of recovery from inactivation on channel stoichiometry that would not require us to discard the view that dissociation of a single inactivation determines recovery? One simple explanation of this result is that it does not reflect some unusual aspect of the inactivation mechanism per se but rather reflects the coupling of recovery from inactivation to Ca$^{2+}$-dependent activation steps. In fact, for both BK$\alpha$ currents in chromaffin cells and $\alpha + \beta_2$ currents, the time course of recovery from inactivation becomes faster both at more negative potentials and with reductions in cytosolic Ca$^{2+}$ (Ding and Lingle, 2002). Thus, although dissociation of a single inactivation particle may be sufficient to remove inactivation, it seems likely that, dependent on Ca$^{2+}$ and recovery potential, channels may reinactivate during the recovery process. As a consequence, the time course of recovery from inactivation most likely reflects multiple kinetic steps, including dissociation of an inactivation domain, but also other Ca$^{2+}$-dependent transitions. Because reductions in the $\beta_2\alpha$ injection ratio shift the $V_{0.5}$ for activation to more positive values, this would naturally then be expected to also produce effects on recovery from inactivation.

Two sets of experiments were done to test this idea. First, we examined the time course of recovery from inactivation after different amounts of removal of inactivation of $\alpha + \beta_2$ currents by trypsin. Similar to our previous results (Ding et al., 1998; Li et al., 1999), when the average number of inactivation domains per channel is altered by digestion with trypsin, the time course of recovery from inactivation remains virtually unchanged (results not shown). The difference between the experiment with trypsin and the results with $\beta_2\alpha$ dilution is that, in the former case, channels with fewer inactivation domains still have a full set of $\beta$ subunits per channel, thereby leaving the voltage dependence of activation unchanged. In a second type of experiment, the average number of inactivation domains per channel was altered not by dilution of $\beta_2$ subunits but by coexpression of $\beta_2$ and $\alpha$ subunits, along with a $\beta_2$ subunit in which the N-terminal inactivation domain has been removed (construct $\beta_2$-Δ33; Xia et al., 1999). Thus, a full complement of $\beta_2$ subunits will be available to associate with $\alpha$ subunits, but less than a full complement of inactivation domains will be present. We varied the ratio of

![Figure 8. Recovery from inactivation shifts with changes in the $\beta_2\alpha$ subunit ratio. A. Currents were activated with 10 μm with the paired pulse recovery protocol shown on the top. Each set of traces corresponds to a different $\beta_2\alpha$ injection ratio as indicated. Note the much slower recovery from inactivation at 1.0 $\beta_2\alpha$ than at lower ratios. The dashed lines indicate the 0-current level. B. The percent recovery ± SD is displayed as a function of recovery interval for a set of four to six patches at each injection ratio. The recovery time points at a ratio of 1.0 are obscured by those at 2.0. The recovery time course was fit in each case with a single exponential. For ratios of 0.01, 0.25, 0.05, 0.1, 1.0, and 2.0, $\tau_r$ was 3.9, 4.8, 6.5, 9.0, 19.3, and 19.0 msec, respectively. C. The dependence of $\tau_r$ on the injection ratio is displayed.](image-url)
functions of both inactivation and activation transitions, mechanistic interpretation of the correspondence of these parameters with β2 subunit stoichiometry is substantially more complicated. Thus, we present these figures primarily for their physiological significance, i.e., that alteration of β2:α subunit stoichiometry can affect key functional properties of the resulting α + β2 currents that are likely to influence the role such channels can play among different cells.

Each β1 subunit also produces an incremental effect on shifts in V_{0.5} at a given Ca^{2+}

Previous work suggested that a single β1 subunit in association with four α subunits might be sufficient to produce a full shift in the V_{0.5} for activation (Jones et al., 1999). Therefore, a few experiments were done to test whether the observations described above were unique to the β2 subunit. Currents resulting from activation of α + β1 channels were studied after injection of different ratios of β1:α subunit (Fig. 10A). These currents resulted in G–V curves that exhibited a relatively parallel shift (Fig. 10B) more similar to the predictions of the incremental model for β subunit action than an all-or-none model. We also examined G–V curves at injection ratios likely to result in intermediate stoichiometries (e.g., 0.1:1:α) at different days after injection to try to identify conditions that might mimic previous results (Jones et al., 1999). In no case have we observed G–V curves that exhibit the clear separation into distinct Boltzmann components that was observed previously (Jones et al., 1999).

To verify that a range of stoichiometric combinations of β1 and α subunits were occurring, we also examined the activation characteristics of β1 channels in which the N terminus from the β2 subunit was appended (construct β1–C2). Coexpression of α + β1–C2 subunits resulted in inactivating currents similar to wild-type α + β2 currents (Fig. 10C), although the limiting τ appeared a bit slower. If the assembly of α + β1 subunits was such that only 0.4 and 4:4 combinations were occurring, we also examined the activation characteristics of β1 channels in which the N terminus from the β2 subunit was appended (construct β1–C2). Consistent with this, the G–V curves for activation of α + β1–C2 currents shifted the β1–C2:α subunit ratio in an approximately parallel manner (Fig. 10D).

We therefore evaluated the relationship between τ_i, f_{on}, and V_{0.5} for activation for the α + β1–C2 currents. As shown in Figure 11A, τ_i varies with f_{on} in a manner consistent with an inactivation mechanism similar to that for the α + β2 currents. In this case, the α + β1–C2 currents are best approximated by a model in which the limiting minimal τ_i is ~30 msec. In Figure 11B, the activation V_{0.5} is plotted as a function of τ_i for both the β1–C2 construct and β2. The relationships are similar, although the V_{0.5} for activation at 10 μM Ca^{2+} is shifted to somewhat more negative potentials for the β1–C2 construct. Consistent with this, the full range of shift in V_{0.5} produced by the β1–C2 construct relative to α alone was closer to ~100 mV rather than the near ~80 mV observed for the β2 construct. On the basis of the analysis described above, the values of τ_i obtained for the β1–C2 construct were then related to the predicted average number of β subunits per channel assuming a τ_{min} of 30 msec and a τ_{max} of 120 msec. Over a broad range of predicted channel stoichiometries, V_{0.5} varies approximately linearly with the predicted number of β subunits per channel (Fig. 11C). This is the result consistent with an incremental effect of each β1 subunit on V_{0.5}}

β2:β2-Δ33 to change the number of inactivation domains per channel in the population. Regardless of the ratio of β2:β2-Δ33, the time course of recovery is indistinguishable whether channels inactivate with time constants of 20–30 or 60–70 msec.

The above results therefore support the view that the seemingly anomalous faster rate of recovery from inactivation with smaller β2:α ratios is simply a consequence of the influence of current activation transitions on the recovery time course (Ding and Lingle, 2002). In essence, τ_i is not dependent on the number of inactivation domains per channel but rather on the number of β subunits per channel because of coupling of the recovery process to steps in the activation pathway. An interesting implication of this result is that variation in τ_i based on the stoichiometry of β2:α subunit assembly suggests a novel mechanism by which key functional properties of BK currents might be regulated.

To summarize how steady-state inactivation and τ_i vary, in accordance with the stoichiometry of those channels, we have plotted V_{0.5} for steady-state inactivation and τ_i as a function of τ_i (Fig. 9A,B). To evaluate the extent to which stoichiometry may affect the inactivation V_{0.5} and τ_i, these parameters were also plotted (Fig. 9C,D) against the predicted average number of β2 subunits per channel based on the analysis of τ_i presented in Figure 4. Because both the inactivation V_{0.5} and τ_i are complex

**Figure 9.** Dependence of steady-state inactivation and recovery from inactivation on β2:α stoichiometry in α + β2 channels. A, Inactivation V_{0.5} determined exclusively for the inactivating portion of current is plotted as a function of the inactivation time constant for the same set of patches. B, τ_i is plotted as a function of the inactivation time constant. C, Inactivation V_{0.5} determined either from the inactivating portion of the current (●) or from the whole channel population (○) is related to the average number of β2 subunits per channel based on the inactivation properties of the currents. The inactivating portion of the current appears to vary with subunit stoichiometry in a manner qualitatively similar and parallel to the activation V_{0.5} (Fig. 4). D, τ_i is plotted as a function of subunit stoichiometry and exhibits a surprisingly steep dependence on the apparent number of β2 subunits per channel. Other experiments described in Results indicate that this does not reflect some complexity in the stoichiometry of the unblocking portion of the inactivation process but rather reflects the coupling of recovery to shifts in the voltage dependence of activation.
channels, the auxiliary subunit appended upstream of the fi
cation activity (McManus et al., 1995; Meera et al., 1996; coassemble with the pore-forming complexes in which one or more kinds of auxiliary subunits exist as representatives of both voltage- and transmitter-gated families, exist as well in virtually all ion channels, including representatives of both voltage- and transmitter-gated families, exist as essentially parallel changes with some reduction in the slope at the intermediate dilutions. For ratios of 0, 0.01, 0.02, 0.1, and 2.0, \( V_{0.5} \) values were +48.9, +28.8, +4.1, −11.4, and −34.9 mV, respectively. C, α subunits were coexpressed with a β1 construct with the β2 N terminus appended upstream of the first transmembrane segment (termed β1–C2). Currents were activated with 10 μM Ca\(^{2+}\) with the voltage protocol shown in A. The presence of inactivation in this construct provides an independent measure of the stoichiometry of assembly. D, G–V curves were generated for a set of patches from traces similar to those shown in C. Activation \( V_{0.5} \) values were +48.9, +17.0, −3.7, and −47.8 mV for ratios of 0, 0.05, 0.1, and 2.0, respectively.

and is inconsistent with the all-or-none effect of a single β1 subunit.

**DISCUSSION**

The pore-forming core of virtually all ion channels consists of a multimeric structure, arising either from assembly of identical or related subunits or from repeated elements contained within the same protein. In addition, many ion channels, including representatives of both voltage- and transmitter-gated families, exist as complexes in which one or more kinds of auxiliary subunits coassemble with the pore-forming α subunits. In the case of BK channels, the auxiliary β subunit KCNMB family appears to play a critical role in defining almost every important functional property of the channel complex, including the apparent Ca\(^{2+}\) dependence of gating (McManus et al., 1995; Meera et al., 1996; Wallner et al., 1999; Xia et al., 1999), activation and deactivation behavior (Brenner et al., 2000), inactivation (Wallner et al., 1999; Xia et al., 1999, 2000; Uebele et al., 2000), and even instantaneous current–voltage behavior (Zeng et al., 2001). Because the stoichiometry of assembly of α and β subunits is 1:1 (Knaus et al., 1994a), the tetrameric BK channels can contain up to four β subunits.

Here we have taken advantage of the properties of inactivation conferred by β2 auxiliary subunits to examine the consequences
of different stoichiometric combinations of $\alpha + \beta_2$ subunits on functional properties of the channels. We first establish that the inactivation properties of macroscopic $\alpha + \beta_2$ currents resulting from different $\beta_2:\alpha$ ratios are generally consistent with the idea that functional channels are formed with less than a full complement of $\beta_2$ subunits. This is also directly supported by single-channel recordings. Using inactivation as an indicator of the average number of $\beta_2$ subunits per channel, we then relate other channel functional properties to the channel stoichiometry. The comparison of both macroscopic and single-channel currents argues that each $\beta_2$ subunit contributes incrementally to influence the activation $V_{0.5}$ at a given $[\text{Ca}^{2+}]$. A similar conclusion was obtained with the $\beta_1$ subunit. In addition to activation $V_{0.5}$, both the $V_{0.5}$ for steady-state inactivation and the time constant of recovery from inactivation varied with channel stoichiometry. The pronounced effect of $\beta_2:\alpha$ stoichiometry on inactivation $V_{0.5}$ and $\tau_s$ supports the view that variation in BK channel properties in cells can be continuously varied by the ratio of $\beta$ to $\alpha$ subunits.

**Dependence of gating behavior on $\beta_2:\alpha$ stoichiometry**

Compared with channels containing only $\alpha$ subunits, the $\beta_2$ subunit shares with the $\beta_1$ subunit an ability to shift the equilibrium between closed and open states to more negative voltages at a given $[\text{Ca}^{2+}]$. The present results suggest that association of each $\beta_2$ subunit independently exerts an incremental effect on the gating equilibrium. On the basis of inactivation time constants, we were able to make inferences about the average number of $\beta_2$ subunits per channel from different combinations, thereby resulting in the functional assembly of inactivating and noninactivating subunits (Ding et al., 1998). However, how this assembly might relate to activation $V_{0.5}$ was not addressed. In human coronary smooth muscle, BK channels were observed to exhibit five different $P(0)$ versus $V$ curves, incrementally spanning over 120 mV (Tanaka et al., 1997), presumably reflecting five populations of possible $\beta_1:\alpha$ subunit assemblies. Similarly, BK channels from the avian nasal salt gland, when studied in lipid bilayers, also clustered into five different apparent $\text{Ca}^{2+}$ sensitivities (Wu et al., 1996), which were interpreted to reflect the five stoichiometric combinations of two subunits, one of higher and one of lower $\text{Ca}^{2+}$ affinity.

These previous studies contrast with the conclusions of one other study in which it was suggested that the contribution of a single $\beta_1$ subunit to a channel was sufficient to produce an all-or-none effect on the shift in gating (Jones et al., 1999). The main points in favor of the all-or-none model were the following. As the ratio of $\beta_1:\alpha$ increased, the shift in activation $V_{0.5}$ for macroscopic currents exhibited a very steep change from $\alpha$- to $\alpha + \beta_1$-like. For those patches with a $V_{0.5}$ intermediate between the two extremes, the $G-V$ curves exhibited two-component Boltzmann relationships similar to the predictions of the all-or-none model (Fig. 3D). Furthermore, when kinetic properties of single channels in patches from oocytes injected with low $\beta_1:\alpha$ ratios were examined, individual channels exhibited either of two different open interval behaviors, one comparable with Slo1 channels alone and one with a longer open time. We consider these results quite convincing that channels with a behavior intermediate between that with $\alpha$ alone and that with $4\beta_1:4\alpha$ were not observed, although we know of no simple way to reconcile these previous results with our own.

The present results have the advantage that the inactivation properties provide an independent indication that channels with intermediate stoichiometries do actually occur. Thus, one possibility is that the conditions of expression in the Jones study somehow resulted in channels with either 0:4 or 4:4 $\beta_1:\alpha$ stoichiometries, although it is difficult to imagine how this might occur. One difference between the study of Jones et al. (1999) and our own is that the molar $\beta_1:\alpha$ ratios used here appear substantially higher than those used in the other study. Another difference was the use of 4:26 and 3:31 $\alpha$ subunit splice variants (Jones et al., 1999) in contrast to the 0:3 variant used here.

Although these explanations remain unsatisfying, for the present we propose that, when a direct measure of subunit stoichiometry is available, each $\beta$ subunit produces an incremental effect on channel gating. However, under some as yet undefined conditions, $\beta_2$ subunit assembly is sufficiently highly cooperative that the resulting channel population can contain primarily 0:4 and 4:4 $\beta_2$ combinations, thereby resulting in the functional equivalent of an all-or-none $\beta$ subunit effect.

**$\beta$ subunits and the functional diversity of BK channel properties**

BK channels in native tissues exhibit great diversity in their functional properties (McManus, 1991; Vergara et al., 1998), and, most notably, different BK channels exhibit substantial variation in their activation ranges at a given $[\text{Ca}^{2+}]$. This reflects an important contribution of different $\beta$ subunits to the BK channel complex among different tissues, with some lesser contribution of different $\alpha$ subunit splice variants.

Variation in the average stoichiometry of $\beta_2:\alpha$ subunits among cells clearly can play a key role in defining the gating properties of the BK channels. Such a mechanism has been proposed to ac-
count for the role of BK channels in frequency tuning in hair cells (Jones et al., 1998, 1999). The present results provide additional support for the idea that the activation range for a population of BK channels can be continuously adjusted on the basis of the fractional occupancy of the each α subunit with an appropriate β subunit. Furthermore, for inactivating BK channels, both channel availability and the rate of return from inactivation also vary continuously as a function of the average βα subunit stoichiometry within the channel population. Thus, these results indicate that essentially every physiologically important parameter of BK channel function can be substantially regulated by adjustment of the relative expression of β to α subunits.

REFERENCES


