

# The $\beta 1$ Subunit Enhances Oxidative Regulation of Large-Conductance Calcium-activated $K^+$ Channels

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**ABSTRACT** Oxidative stress may alter the functions of many proteins including the Slo1 large conductance calcium-activated potassium channel ( $BK_{Ca}$ ). Previous results demonstrated that in the virtual absence of  $Ca^{2+}$ , the oxidant chloramine-T (Ch-T), without the involvement of cysteine oxidation, increases the open probability and slows the deactivation of  $BK_{Ca}$  channels formed by human Slo1 (*hSlo1*)  $\alpha$  subunits alone. Because native  $BK_{Ca}$  channel complexes may include the auxiliary subunit  $\beta 1$ , we investigated whether  $\beta 1$  influences the oxidative regulation of *hSlo1*. Oxidation by Ch-T with  $\beta 1$  present shifted the half-activation voltage much further in the hyperpolarizing direction ( $-75$  mV) as compared with that with  $\alpha$  alone ( $-30$  mV). This shift was eliminated in the presence of high  $[Ca^{2+}]_i$ , but the increase in open probability in the virtual absence of  $Ca^{2+}$  remained significant at physiologically relevant voltages. Furthermore, the slowing of channel deactivation after oxidation was even more dramatic in the presence of  $\beta 1$ . Oxidation of cysteine and methionine residues within  $\beta 1$  was not involved in these potentiated effects because expression of mutant  $\beta 1$  subunits lacking cysteine or methionine residues produced results similar to those with wild-type  $\beta 1$ . Unlike the results with  $\alpha$  alone, oxidation by Ch-T caused a significant acceleration of channel activation only when  $\beta 1$  was present. The  $\beta 1$  M177 mutation disrupted normal channel activation and prevented the Ch-T-induced acceleration of activation. Overall, the functional effects of oxidation of the *hSlo1* pore-forming  $\alpha$  subunit are greatly amplified by the presence of  $\beta 1$ , which leads to the additional increase in channel open probability and the slowing of deactivation. Furthermore, M177 within  $\beta 1$  is a critical structural determinant of channel activation and oxidative sensitivity. Together, the oxidized  $BK_{Ca}$  channel complex with  $\beta 1$  has a considerable chance of being open within the physiological voltage range even at low  $[Ca^{2+}]_i$ .

**KEY WORDS:**  $BK_{Ca}$  • *hSlo* • chloramine-T • methionine • cysteine

## INTRODUCTION

The large conductance calcium-activated potassium channel ( $BK_{Ca}$ ) exists in various types of cells and tissues including smooth muscle and brain. In response to depolarization and/or a rise in intracellular  $Ca^{2+}$  ( $[Ca^{2+}]_i$ ),  $BK_{Ca}$  channels mediate net  $K^+$  efflux to repolarize the membrane potential to the resting state. This function serves an important role in muscle contraction—during which  $Ca^{2+}$  sparks activate the  $BK_{Ca}$  channels leading to vasorelaxation (Nelson et al., 1995; Jaggar et al., 2000)—and the afterhyperpolarization phase of the action potential in select neurons (Storm, 1987). Furthermore, the impairments exhibited by mice lacking the channel indicate that  $BK_{Ca}$  channels influence normal urinary bladder (Meredith et al., 2004) and cerebellar functions (Sausbier et al., 2004).

The human  $BK_{Ca}$  channel pore-forming  $\alpha$  subunit (*hSlo1*) contains seven putative transmembrane-spanning

regions (Dworetzky et al., 1994; Pallanck and Ganetzky, 1994; Tseng-Crank et al., 1994). The S0 transmembrane domain, which distinguishes the *Slo* from the *Shaker* family of voltage-dependent potassium channels, is thought to be a site of interaction with auxiliary  $\beta$  subunits (Wallner et al., 1996; Meera et al., 1997). Multiple types of  $\beta$  subunits ( $\beta 1$ –4) have been isolated in mammals, each with a different tissue distribution and function (Knaus et al., 1994; Xia et al., 1999; Brenner et al., 2000a; Uebele et al., 2000).

The  $\beta 1$  subunit is a 25-kD membrane protein consisting of two transmembrane domains connected by a large extracellular loop, such that both the  $NH_2$  and  $COOH$  termini are intracellularly located (Knaus et al., 1994; Orio et al., 2002; Patterson et al., 2002). The  $\beta 1$  subunit is present in the brain, particularly in the hippocampus

*Abbreviations used in this paper:*  $BK_{Ca}$  channel, large conductance calcium-activated potassium channel; Ch-T, chloramine-T;  $\Delta G_{Ca}$ , change in free energy change associated with  $Ca^{2+}$  binding; met-O, methionine sulfoxide;  $Q_{app}$ , apparent equivalent charge movement; ROS/RNS, reactive oxygen/nitrogen species;  $V_{0.5}$ , half-activation voltage; z, equivalent charge.

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and corpus callosum (Tseng-Crank et al., 1996), but is predominantly expressed in smooth muscle (Garcia-Calvo et al., 1994; Tanaka et al., 1997). The impaired vasorelaxation found in  $\beta 1$  knockout mice (Brenner et al., 2000b; Pluger et al., 2000) and the down-regulation of  $\beta 1$  expression associated with some forms of hypertension (Gollasch et al., 2002; Amberg et al., 2003; Amberg and Santana, 2003) clearly underscore the important physiological role of  $\beta 1$  in the  $BK_{Ca}$  channel regulation of vascular function. The presence of  $\beta 1$  modulates  $BK_{Ca}$  channel activity by enhancing the apparent  $Ca^{2+}$  sensitivity of the pore-forming subunit and also by slowing the activation/deactivation kinetics, even in the virtual absence of  $Ca^{2+}$  (McManus et al., 1995; Wallner et al., 1995; Meera et al., 1996; Nimigeon and Magleby, 1999, 2000; Cox and Aldrich, 2000; Qian and Magleby, 2003). The structural determinants within  $\beta 1$  responsible for these critical modulatory properties are just beginning to be identified (Fernandez-Fernandez et al., 2004).

Other regulatory mechanisms such as phosphorylation, pH, and the cellular redox state influence  $BK_{Ca}$  channel activity (Weiger et al., 2002). During oxidative stress, cellular reactive oxygen/nitrogen species (ROS/RNS) readily modify cysteine and methionine residues in proteins. Oxidation of cysteine typically leads to the formation of disulfides, whereas oxidation of methionine residues creates the polar methionine sulfoxide (met-O). Oxidative modifications of amino acids differentially influence  $BK_{Ca}$  channel function depending on the ROS/RNS, the residues modified within the channel, as well as the experimental model system (DiChiara and Reinhart, 1997; Sobey et al., 1997; Wang and Wu, 1997; Wang et al., 1997; Barlow et al., 2000; Gong et al., 2000; Soh et al., 2001; Brakemeier et al., 2003). Studies using heterologously expressed *hSlo1* indicate that oxidation of cysteine residues typically decreases the channel open probability (DiChiara and Reinhart, 1997; Soto et al., 2002; Tang et al., 2004). In contrast, methionine oxidation of the *hSlo1* pore-forming subunit that is promoted by the oxidant chloramine-T (Ch-T) increases the channel open probability (Tang et al., 2001).

Oxidative stress is prominently involved in many disease states such as vascular dysfunction (Taniyama and Griendling, 2003) and neurodegenerative diseases (Knight, 1997; Markesbery, 1997; Butterfield et al., 2001). These physiological systems that are affected by oxidative stress depend on  $BK_{Ca}$  channel activity for normal function. Therefore, determining the effect of oxidative modification of  $BK_{Ca}$  channel complexes that closely resemble native channels is important to understand and possibly treat or prevent these diseases. Native  $BK_{Ca}$  channels are often multi-subunit

complexes containing both *Slo1* and auxiliary  $\beta$  subunits (Garcia-Calvo et al., 1994; Knaus et al., 1994; Giangiacoia et al., 1995; Vogalis et al., 1996; Tanaka et al., 1997; Wanner et al., 1999; Weiger et al., 2000). However, the influence of  $\beta$  subunits on the oxidative regulation of *Slo1* function has not been thoroughly examined.

The purpose of the present work was to determine whether the presence of  $\beta 1$  alters the functional effects of *hSlo1* oxidation. Methionine oxidation of *hSlo1* alone causes a shift in the macroscopic G-V curve by  $-30$  mV and slows deactivation without any appreciable effect on the activation kinetics at depolarized voltages (Tang et al., 2001). We show that, in the virtual absence of  $Ca^{2+}$ , the auxiliary subunit  $\beta 1$  dramatically potentiates the effect of methionine oxidation in the *hSlo1* pore-forming protein. This is demonstrated by a further increase in the open probability and even greater slowing of the deactivation kinetics. Furthermore,  $\beta 1$  confers novel oxidation sensitivity to the channel activation kinetics that is mediated largely by a single methionine residue located in the second transmembrane domain (TM2) of  $\beta 1$ .

## MATERIALS AND METHODS

### *Channel Expression and Mutagenesis*

*hSlo1* (U11058, hbr1; Tseng-Crank et al., 1994) channel alone, or *hSlo1* and  $\beta 1$  (1:1 weight ratio) were transiently expressed in HEK-tSA cells using FuGENE 6 (Roche Applied Science) as described previously (Avdonin et al., 2003). The mouse *Slo*  $\beta 1$  (*m* $\beta 1$ ; AF020711; Jiang et al., 1999) in pEGFP-N1 (BD Biosciences) was obtained from the laboratory of R. Aldrich (Stanford University, Stanford, CA). The *m* $\beta 1$  mutants M7L, M23L, M177L, and Triple (M7L:M23L:M177L) were constructed using PCR-based mutagenesis, and the sequences were verified. "Cysless" *b* $\beta 1$ , in which every cysteine in bovine  $\beta 1$  (*b* $\beta 1$ ; L26101; Knaus et al., 1994) was replaced with alanine (C18A, C53A, C76A, C103A, and C135A; Hanner et al., 1998), was obtained from the laboratory of M.L. Garcia (Merck Research Laboratories, Rahway, NJ).

### *Electrophysiology and Data Analysis*

Currents were recorded from excised inside-out patches at room temperature essentially as described previously (Tang et al., 2001). Patch electrodes (Warner) had a typical initial resistance of 2.5–3 M $\Omega$  when filled with solutions (described in the next section); the series resistance,  $\sim 90\%$  of the input resistance, was electronically compensated. The current signal was filtered at 10 kHz through the built-in filter of the patch-clamp amplifier (model AxoPatch 200A; Axon Instruments). Data were acquired and analyzed using Pulse/PulseFit (HEKA), PatchMachine (Avdonin et al., 2003), and IgorPro (WaveMetrics) as described for single-channel data (Avdonin and Hoshi, 2001) and macroscopic current data (Tang et al., 2001; Avdonin et al., 2003). In brief, normalized macroscopic conductance was estimated from single exponential fits to the tail currents recorded at  $-50$  mV excluding the initial 180  $\mu$ s after pulses to different voltages from the holding voltage of 0 mV. The apparent equivalent charge movement ( $Q_{app}$ ) was derived from the simple Boltzmann function

used to describe the average G-V curve. Activation and deactivation time courses were fitted by single exponentials excluding the initial 150- and 180- $\mu$ s segments, respectively. A single exponential fit to the voltage dependence of the time constant provided the value of the equivalent charge movement ( $z$ ).

In some patches, the tail currents after Ch-T treatment contained a minor fast component. The fractional amplitude of this component was typically small (<10%), and the time constant estimated from single-exponential fits was essentially the same as that of the slow component estimated from two-exponential fits. Thus, single-exponential fits were used throughout to quantify the tail current kinetics. Because the time constant of the tail current before modification and that of the minor fast component after Ch-T treatment were similar, the fast component likely reflects the kinetics of unmodified channels.

The change in free energy associated with  $\text{Ca}^{2+}$  binding ( $\Delta G_{\text{Ca}}$ ) was determined based on the  $\Delta G_{\text{Ca}}$  contribution to channel open probability ( $P_o$ ) as described previously (Tang et al., 2004). The values of  $P_o$ ,  $\Delta G_o$ ,  $\Delta G_v$ , and  $\Delta G_{\text{Ca}}$  were estimated by fitting the G-V curves obtained in 0 and 2.1  $\mu\text{M}$   $\text{Ca}^{2+}$ .

Statistical comparisons were made using the paired  $t$  test. In some cases, the  $t$  test and ANOVA followed by the Bonferroni post hoc test were used as specifically indicated (DataDesk; Data Description). Statistical significance was assumed at  $P \leq 0.05$ . Where appropriate, data are presented as mean  $\pm$  SEM.

### Reagents and Solutions

Both the external and internal recording solutions contained the following (mM): 140 KCl, 11 EGTA, and 10 HEPES, pH 7.2 adjusted with NMDG. The free  $\text{Ca}^{2+}$  concentration for these solutions was estimated at <1 nM assuming 20  $\mu\text{M}$  contaminating  $\text{Ca}^{2+}$  (Patcher's Power Tools v1.0, F. Mendez; <http://www.mpibpc.gwdg.de/abteilungen/140/software/>). The external solution used to reduce the size of inward  $\text{K}^+$  currents for experiments involving 2.1  $\mu\text{M}$   $[\text{Ca}^{2+}]_i$  contained the following (mM): 70 KCl, 70 NaCl, 2  $\text{MgCl}_2$ , and 10 HEPES, pH 7.2 adjusted with NMDG. The 2.1- $\mu\text{M}$  free  $\text{Ca}^{2+}$  internal solution contained the following (mM): 120 KCl, 20 KOH, 1  $\text{MgCl}_2$ , 2.2  $\text{CaCl}_2$ , 4 HEDTA, and 10 HEPES, pH 7.4 adjusted with NMDG. The external solution used for experiments involving 120  $\mu\text{M}$   $[\text{Ca}^{2+}]_i$  contained the following reagents (mM): 140 KCl, 2  $\text{MgCl}_2$ , and 10 HEPES, pH 7.2 adjusted with NMDG. The 120- $\mu\text{M}$  free  $\text{Ca}^{2+}$  internal solution contained the following reagents (mM): 140 KCl, 10  $\text{MgCl}_2$ , 0.1  $\text{CaCl}_2$ , and 10 HEPES, pH 7.2 adjusted with NMDG.

Chloramine-T (Ch-T; Sigma-Aldrich) was dissolved in the internal solution immediately before use. In every experiment, 2 mM Ch-T was manually applied with a pipette to ensure the addition of six times the bath volume ( $\sim 150 \mu\text{l}$ ). With Ch-T present, channel current in response to a pulse to 120 mV was monitored every 5 s for the following three features of oxidation by Ch-T: increased current amplitude, slowed deactivation, and accelerated activation. Once these characteristic changes reached steady-state levels ( $\leq 8$  min), Ch-T was subsequently washed out with 1 ml of recording solution. The time courses of modification of channels, composed of either *hSlo1* alone or *hSlo1* and  $\beta 1$  together, were indistinguishable.

## RESULTS

### Oxidation by Ch-T More Dramatically Enhances *hSlo1* Currents When $\beta 1$ Is Present

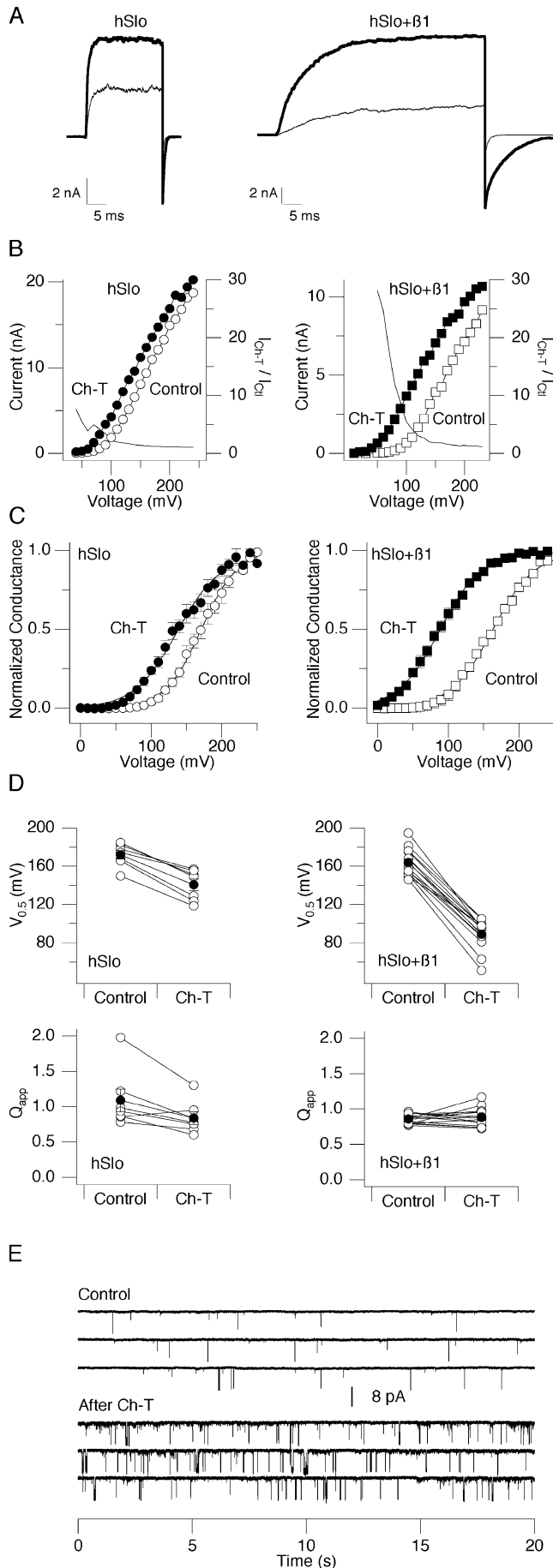
To determine if the presence of  $\beta 1$  influences the functional effects of *hSlo1* oxidation by Ch-T, ionic currents through *hSlo1* or *hSlo1* +  $\beta 1$  channels were recorded

in the inside-out patch-clamp configuration from transiently transfected HEK-tsA cells. All recordings were initially made in the virtual absence of  $\text{Ca}^{2+}$ , essentially permitting the Slo channel to act as a voltage-dependent channel to simplify the data analysis (Meera et al., 1996; Horrigan and Aldrich, 1999; Horrigan et al., 1999). Currents elicited by pulses to 120 mV in patches containing either *hSlo1* or *hSlo1* +  $m\beta 1$  are shown in Fig. 1 A (thin sweeps). The *hSlo1* +  $m\beta 1$  currents displayed slow activation and deactivation (also see Fig. 2), a hallmark of the functional presence of the  $\beta 1$  subunit. After bath application of 2 mM Ch-T to the cytoplasmic side, *hSlo1* and *hSlo1* +  $m\beta 1$  exhibited similar modification time courses ( $P = 0.08$ ,  $t$  test) that resulted in larger current amplitudes (Fig. 1 A, thick sweeps). The current enhancement remained after Ch-T washout, consistent with the oxidative modification of the channel protein complex by Ch-T.

Treatment with Ch-T shifted the peak I-V curves from both *hSlo1* and *hSlo1* +  $m\beta 1$  to more negative voltages, such that at a given voltage, the current size was greater (Fig. 1 B). However, the current enhancement was drastically larger in *hSlo1* +  $m\beta 1$  than in *hSlo1* alone, especially at moderately depolarizing voltages (50–100 mV; Fig. 1 B). The relative increase in current amplitude due to oxidation became progressively smaller at more depolarizing voltages where the channel open probability is saturated. This voltage dependence is consistent with Ch-T increasing the open channel probability as shown for *hSlo1* (Tang et al., 2001).

The voltage dependence of the probability of the channel being open inferred from normalized macroscopic G-V curves confirmed that treatment with Ch-T enhanced the open probability of *hSlo1* +  $m\beta 1$  more profoundly than that of *hSlo1* alone. The G-V curves estimated from tail current measurements were fit by a simple Boltzmann function as a data descriptor to describe the overall voltage dependence of the Ch-T effect (Fig. 1 C). After Ch-T treatment, the *hSlo1* half-activation voltage ( $V_{0.5}$ ) shifted by  $\sim 30$  mV in the hyperpolarizing direction. However, for *hSlo1* +  $m\beta 1$ , oxidation by Ch-T produced the strikingly greater shift of  $-75$  mV. The mean shift in  $V_{0.5}$  for *hSlo1* +  $m\beta 1$  ( $\Delta V_{0.5} = -74.6 \pm 3.5$  mV,  $n = 14$ ) was more than twice as great as  $\Delta V_{0.5}$  for *hSlo1* alone ( $\Delta V_{0.5} = -31.3 \pm 3.3$  mV,  $n = 7$ ) (Fig. 1 D;  $P < 0.0001$ ,  $t$  test). These results suggest that treatment with Ch-T leads to an increase in the open probability that is markedly potentiated with  $m\beta 1$  present.

The apparent equivalent charge movement ( $Q_{\text{app}}$ ) of *hSlo1* activation, inferred from the steepness of the G-V curve, decreased by  $\sim 23\%$  ( $\Delta Q_{\text{app}} = -0.25 \pm 0.07e$ ,  $P = 0.036$ ,  $n = 7$ ) after Ch-T treatment (Fig. 1 D). In contrast, the  $\Delta Q_{\text{app}}$  for *hSlo1* +  $m\beta 1$  demonstrated no significant change after modification ( $\Delta Q_{\text{app}} = 0.02 \pm 0.02e$ ;  $P = 0.65$ ,  $n = 14$ ).

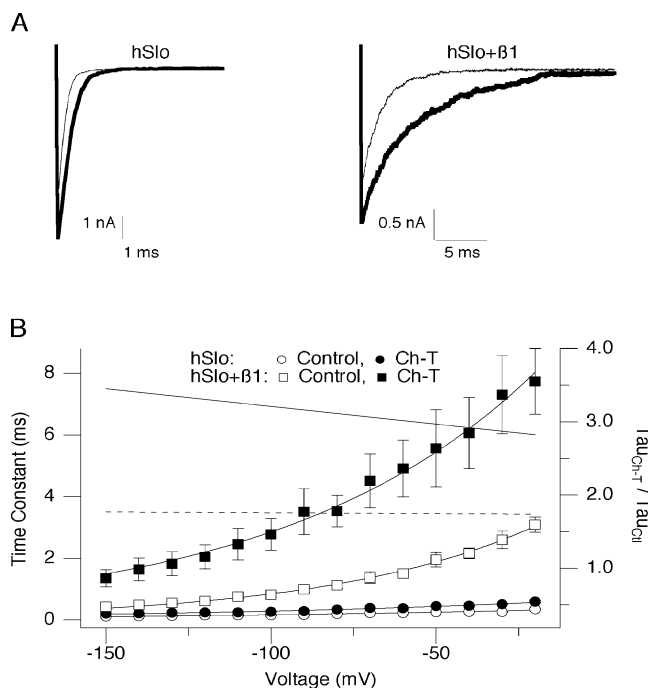


The increase in the channel open probability caused by Ch-T was maintained at more negative, physiological voltages. At  $-40$  mV in the virtual absence of  $Ca^{2+}$ , treatment with Ch-T markedly increased the number of hSlo1 + mβ1 channel openings (Fig. 1 E). Indeed, the mean open probability at this voltage increased by a factor of  $12.0 \pm 3.0$  relative to control. In contrast with the dramatic changes in the gating properties of the hSlo1 + mβ1 channel, the open channel current-conductance characteristic estimated using voltage ramps (0–250 mV) in single-channel patches remained unaltered by Ch-T treatment (unpublished data).

#### Modification by Ch-T Drastically Slows hSlo1 + mβ1 Deactivation

To assess whether treatment with Ch-T affects hSlo1 deactivation differently when the β1 subunit is present, hSlo1 and hSlo1 + mβ1 tail currents were recorded before and after Ch-T treatment (Fig. 2 A). After Ch-T exposure, the mean deactivation time constant at  $-40$  mV increased by  $\sim 70\%$  (from 0.26 to 0.45 ms) for hSlo1, whereas the increase for hSlo1 + mβ1 was  $\sim 180\%$  (from 2.12 to 6.06 ms). This appreciably greater slowing of hSlo1 + mβ1 deactivation was observed at every voltage examined (Fig. 2 B). Single exponential fits to the voltage dependence of the deactivation time constants in the voltage range of  $-150$  to  $-50$  mV indicated that oxidation by Ch-T specifically increased the time constant values at 0 mV,  $\tau(0)$ , for hSlo1 and hSlo1 + mβ1 ( $P = 0.0021$  and  $0.015$ , respec-

FIGURE 1. Oxidation by Ch-T enhances hSlo1 + mβ1 currents to a greater extent than hSlo1 currents. (A) Representative currents before (thin sweep) and after (thick sweep) 2 mM Ch-T treatment. The currents were elicited in response to pulses from 0 to 120 mV. Mean times to reach 50% of final current amplitude in the presence of Ch-T for hSlo1 and hSlo1 + mβ1 were  $5.64 \pm 0.34$  min and  $4.68 \pm 0.3$  min, respectively ( $P = 0.08$ ,  $n = 4$ ). (B) Peak I-V curves before (open symbols) and after (closed symbols) modification by Ch-T. Continuous curves represent relative increases in current amplitude as a function of voltage (right axis). (C) G-V curves before (open symbols) and after (closed symbols) modification by Ch-T. The macroscopic currents were elicited by pulses to different test voltages from the holding voltage of 0 mV. The hSlo1  $V_{0.5}$  values for the results obtained before and after Ch-T application were  $171.9 \pm 4.5$  mV and  $140.6 \pm 6.1$  mV ( $\Delta V_{0.5}$  range,  $-19$  to  $-42$  mV;  $P < 0.0001$ ,  $n = 7$ ), respectively. The hSlo1 + mβ1  $V_{0.5}$  values for the results obtained before and after Ch-T application were  $163.8 \pm 3.8$  mV and  $89.2 \pm 4.1$  mV ( $\Delta V_{0.5}$  range  $-50$  to  $-99$  mV;  $P < 0.0001$ ,  $n = 14$ ), respectively. The hSlo1  $Q_{app}$  values for the results obtained before and after Ch-T application were  $1.09 \pm 0.16e$  and  $0.84 \pm 0.09e$  ( $P = 0.036$ ,  $n = 7$ ), respectively. The hSlo1 + mβ1  $Q_{app}$  values for the results obtained before and after Ch-T application were  $0.86 \pm 0.02e$  and  $0.88 \pm 0.04e$ , respectively, ( $P = 0.65$ ,  $n = 14$ ). (D)  $V_{0.5}$  and  $Q_{app}$  values before and after oxidation by Ch-T from individual experiments (open circles) and mean values (closed circles). (E) Representative hSlo1 + mβ1 channel openings at  $-40$  mV before and after treatment with Ch-T. Data were filtered at 10 kHz and sampled at 83 kHz, but are shown filtered at 1 kHz for display purpose. Typically, 3–11 segments were analyzed in each condition.

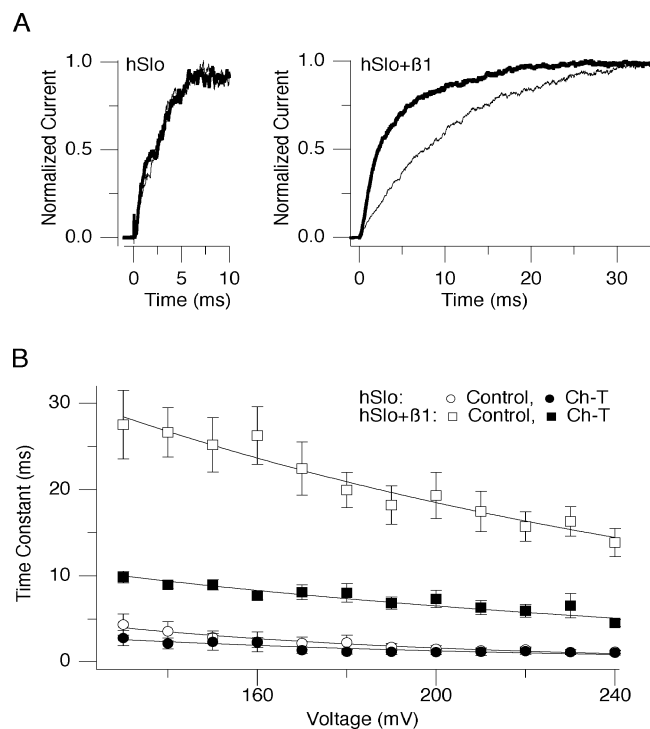


**FIGURE 2.** Ch-T treatment slows deactivation of *hSlo1* + *mβ1* to a greater extent than *hSlo1* deactivation. (A) Tail currents recorded at  $-40$  mV after pulses to  $180$  mV before (thin sweep) and after (thick sweep) Ch-T treatment. (B) Voltage dependence of the deactivation time constant for *hSlo1* control (open circles;  $n = 7$ ), *hSlo1* after Ch-T (closed circles;  $n = 7$ ), *hSlo1* + *mβ1* control (open squares;  $n = 5$ ), and *hSlo1* + *mβ1* after Ch-T (closed squares;  $n = 5$ ). The *hSlo1*  $\tau(0)$  and  $z$  values obtained before and after Ch-T application were  $0.35 \pm 0.04$  ms and  $0.19 \pm 0.01e$ , and  $0.63 \pm 0.06$  ms and  $0.21 \pm 0.01e$ , respectively. The *hSlo1* + *mβ1*  $\tau(0)$  and  $z$  values obtained before and after Ch-T application were  $3.96 \pm 0.52$  ms and  $0.38 \pm 0.02e$ , and  $10.9 \pm 2.1$  ms and  $0.34 \pm 0.01e$ , respectively. The relative increase in the value of the deactivation time constant as a function of voltage (right axis) is shown for *hSlo1* (dashed line) and *hSlo1* + *mβ1* (continuous line).

tively,  $n = 5$ ) without significantly affecting their equivalent charge movement ( $P = 0.17$  and  $0.08$ , respectively,  $n = 5$ ). In fact, the change in  $\tau(0)$  for *hSlo1* + *mβ1* is approximated by a voltage shift of  $-75$  mV, which is similar in value to the voltage shift of the G-V curve after oxidation by Ch-T.

#### Activation Kinetics of *hSlo1* + *mβ1* Accelerates after Ch-T Treatment

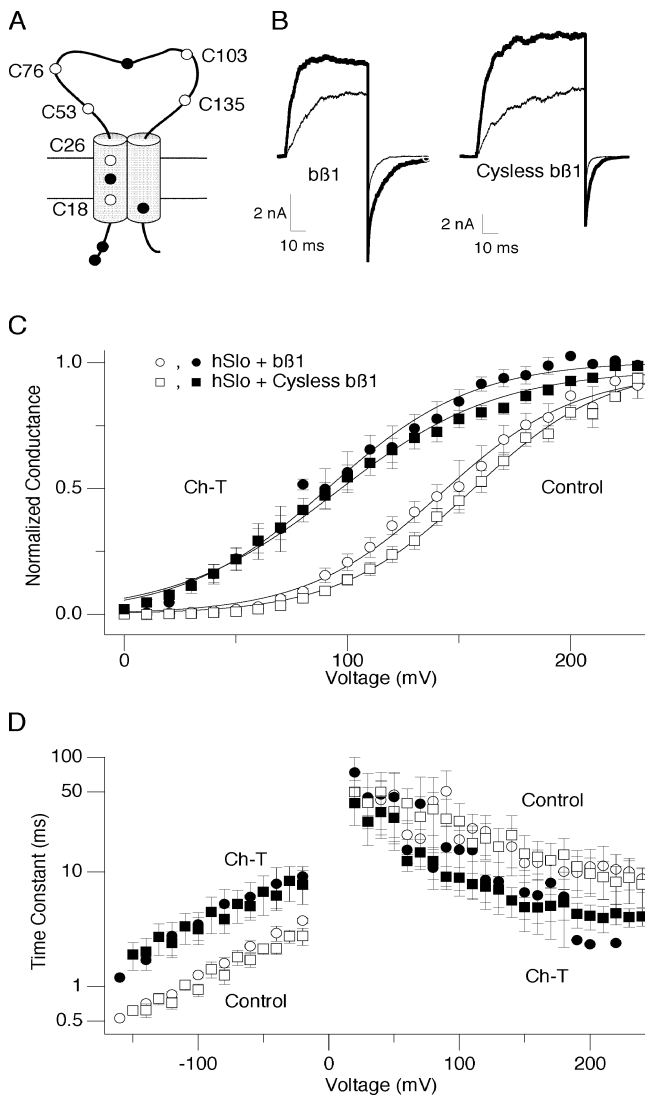
The activation kinetics of *hSlo1* alone remained unaltered after modification by Ch-T (Fig. 3 A; Tang et al., 2001). In contrast, Ch-T markedly accelerated the mean activation kinetics of *hSlo1* + *mβ1* by 68% at each voltage tested (130–240 mV; Fig. 3 B). Single exponential fits to the voltage-dependence of the activation time constant within this voltage range demonstrated that treatment with Ch-T decreased  $\tau(0)$  for *hSlo1* + *mβ1* ( $P = 0.002$ ,  $n = 7$ ), but not for *hSlo1* ( $P = 0.23$ ,  $n = 4$ ), without affecting the equivalent charge movement (*hSlo1*:  $P = 0.14$ ,  $n = 4$ ; *hSlo1* + *mβ1*:  $P = 0.92$ ,  $n = 7$ ).



**FIGURE 3.** Modification by Ch-T accelerates activation of *hSlo1* + *mβ1*. (A) Normalized currents recorded at  $240$  mV from the holding voltage of  $0$  mV before (thin sweep) and after (thick sweep) Ch-T treatment. (B) Voltage dependence of the activation time constant for *hSlo1* control (open circles;  $n = 7$ ), *hSlo1* after Ch-T (closed circles;  $n = 7$ ), *hSlo1* + *mβ1* control (open squares;  $n = 14$ ), and *hSlo1* + *mβ1* after Ch-T (closed squares;  $5 \leq n \leq 12$ ). The *hSlo1*  $\tau(0)$  and  $z$  values obtained before and after Ch-T application were  $7.0 \pm 1.2$  ms and  $0.28 \pm 0.05e$ , and  $4.4 \pm 1.0$  ms and  $0.22 \pm 0.02e$ , respectively. The *hSlo1* + *mβ1*  $\tau(0)$  and  $z$  values obtained before and after Ch-T application were  $0.082 \pm 0.01$  s and  $0.171 \pm 0.02e$ , and  $0.026 \pm 0.003$  s and  $0.174 \pm 0.02e$ , respectively.

This change in  $\tau(0)$  could be accounted for by a voltage shift of more than  $-150$  mV, which is much larger in value than the voltage shift of the G-V curve after oxidation by Ch-T.

Altogether, modification by Ch-T caused a much greater increase in *hSlo1* open probability, an enhanced slowing of deactivation, and a distinct acceleration of activation kinetics when  $\beta 1$  was coexpressed. These Ch-T-induced changes specific to *hSlo1* + *mβ1* may involve any of the following possible mechanisms. First, given that cysteine residues are also potential targets of Ch-T under physiological conditions, oxidation of cysteine within  $\beta 1$  may account for the enhanced oxidative regulation of *hSlo1* + *mβ1*. Second, oxidation of methionine within  $\beta 1$  may synergistically enhance the functional effects of *hSlo1* oxidation. Third, the mere presence of  $\beta 1$  may potentiate the functional outcome of oxidation within the *hSlo1* pore-forming subunit. These possible mechanisms are addressed in the next sections.



**FIGURE 4.** Cysless  $\beta 1$  resembles wild-type  $\beta 1$ . (A) A schematic representation of cysteine residues (open circles) in  $m\beta 1$ . C26 does not exist in  $b\beta 1$ . Closed circles represent methionine residues. (B) Representative currents before (thin sweep) and after (thick sweep) Ch-T treatment. The currents were elicited in response to pulses from 0 to 140 mV. (C) G-V curves before and after modification by Ch-T. The  $hSlo1 + b\beta 1$   $V_{0.5}$  values for the results obtained before (open circles) and after (closed circles) Ch-T application were  $145.5 \pm 8.2$  mV and  $91.4 \pm 9.6$  mV ( $\Delta V_{0.5}$  range,  $-51$  to  $-56$  mV;  $P = 0.0009$ ;  $n = 3$ ), respectively. The  $hSlo1 + Cysless b\beta 1$   $V_{0.5}$  values for the results obtained before (open squares) and after (closed squares) Ch-T application were  $156.6 \pm 4.5$  mV and  $98.5 \pm 7.4$  mV ( $\Delta V_{0.5}$  range,  $-39$  to  $-77$  mV;  $P < 0.0001$ ,  $n = 10$ ), respectively. The  $hSlo1 + b\beta 1$   $Q_{app}$  values for the results obtained before and after Ch-T application were  $0.82 \pm 0.06e$  and  $0.8 \pm 0.04e$  ( $P = 0.8$ ,  $n = 3$ ), respectively. The  $hSlo1 + Cysless b\beta 1$   $Q_{app}$  values for the results obtained before and after Ch-T application were  $0.88 \pm 0.03e$  and  $0.74 \pm 0.05e$ , respectively ( $P = 0.009$ ,  $n = 10$ ). (D) Voltage dependence of the deactivation and activation time constants. Symbols are the same as in C.

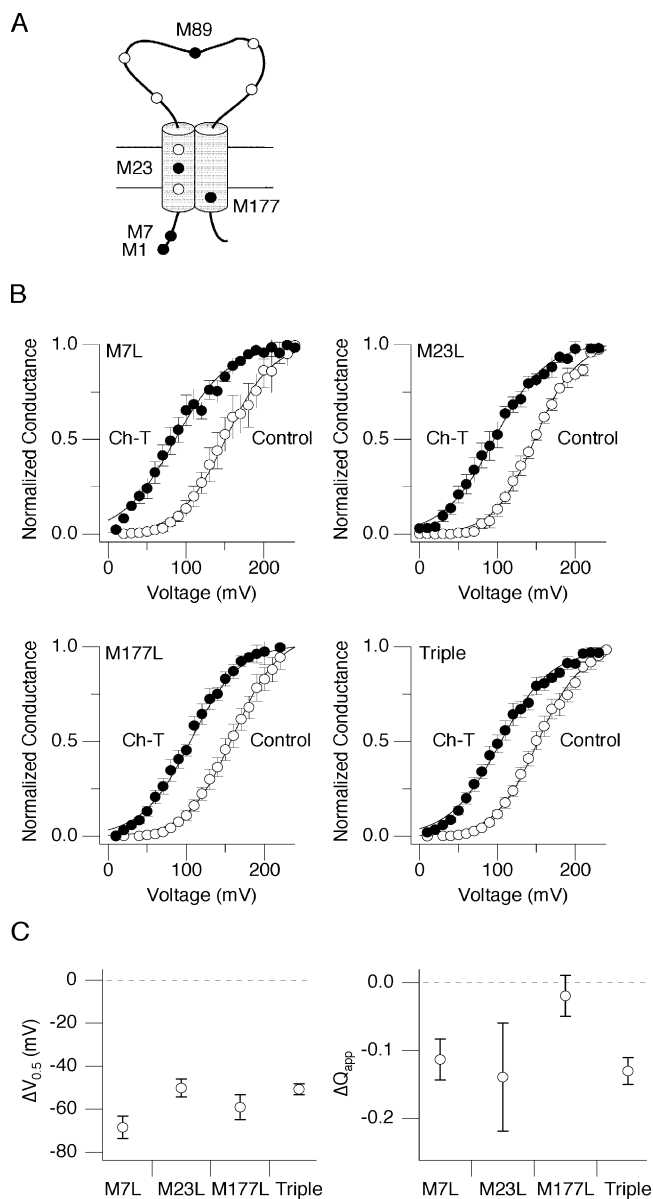
### Cysless $\beta 1$ Produces Results Similar to Wild-type $\beta 1$

Previous results demonstrated that cysteine modification within  $hSlo1$  is not involved in the Ch-T-mediated response (Tang et al., 2001). To test whether the enhanced effects of Ch-T on channel behavior in the presence of  $\beta 1$  involve modification of cysteine residues within the  $\beta 1$  subunit, we used a mutant  $b\beta 1$  subunit devoid of any cysteine named Cysless  $b\beta 1$  (Fig. 4 A; Hanner et al., 1998). Because this mutant was derived from bovine  $\beta 1$ , we compared the results from  $hSlo1 + b\beta 1$  with  $hSlo1 + Cysless b\beta 1$ .

Expression of Cysless  $b\beta 1$  slowed the  $hSlo1$  activation and deactivation kinetics in the control condition essentially as observed with wild-type  $b\beta 1$  (Fig. 4 B), confirming that Cysless  $b\beta 1$  functionally interacts with  $hSlo1$ . Ch-T treatment enhanced the currents through both  $hSlo1 + b\beta 1$  and  $hSlo1 + Cysless b\beta 1$  in a similar manner. After modification by Ch-T, the  $hSlo1 + b\beta 1$  and  $hSlo1 + Cysless b\beta 1$  G-V curves shifted leftward (Fig. 4 C), such that the mean  $\Delta V_{0.5}$  values were indistinguishable ( $-54.1 \pm 1.7$  mV and  $-58.1 \pm 3.6$  mV,  $n = 3$  and 10, respectively). Importantly, these  $\Delta V_{0.5}$  values were markedly greater than those found with  $hSlo1$  alone ( $\Delta V_{0.5} \approx -30$  mV; Fig. 1 D) ( $P < 0.05$ ; Bonferroni test). The  $\Delta V_{0.5}$  values of  $hSlo1 + b\beta 1$  and  $hSlo1 + Cysless b\beta 1$  were smaller than that of  $hSlo1 + m\beta 1$  ( $\Delta V_{0.5} \approx -75$  mV; Fig. 1 D) probably because the control  $V_{0.5}$  values before treatment with Ch-T for the  $b\beta 1$  channel complexes ( $\sim 145$  and  $156$  mV, respectively) were already less depolarized than that of  $m\beta 1$  ( $\sim 164$  mV); yet, all  $V_{0.5}$  values after treatment with Ch-T were  $\sim 90$  mV. Nevertheless, the removal of all cysteine residues within the  $\beta 1$  subunit still permitted the enhanced  $\Delta V_{0.5}$  after modification by Ch-T. Furthermore, the activation and deactivation time courses of  $hSlo1 + Cysless b\beta 1$  before and after treatment with the oxidant resembled those of  $hSlo1 + b\beta 1$  (Fig. 4 D). Therefore, the kinetic and G-V results for  $hSlo1 + m\beta 1$ ,  $hSlo1 + b\beta 1$ , and  $hSlo1 + Cysless b\beta 1$  are largely similar and suggest that oxidation of cysteine residues within  $\beta 1$  is not responsible for the enhanced oxidative regulation of  $hSlo1$  in the presence of the  $\beta 1$  subunit.

### The Greater Increase in Open Probability Does Not Depend on Methionine Oxidation within $\beta 1$

The Ch-T effect on  $hSlo1 + \beta 1$  function did not involve cysteine oxidation but the oxidation of methionine residues within  $\beta 1$  may be responsible. Each  $m\beta 1$  contains five methionine residues: M1, M7, M23, M89, and M177 (Fig. 5 A); the contribution of these methionines to the enhanced shift of the G-V curve and further slowing of deactivation, as well as acceleration of the activation kinetics was assessed in the following manner. Because M1 is obligatory for normal  $\beta 1$  syn-



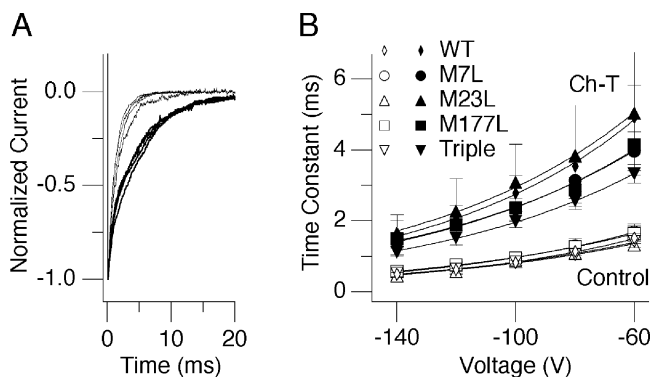
**FIGURE 5.** Methionine mutations within  $m\beta 1$  permit oxidation-related increases in  $hSlo1$  open probability similar to wild-type  $m\beta 1$ . (A) A schematic representation of methionine residues (closed circles) in  $m\beta 1$ . Open circles represent cysteine residues. (B) G-V curves before and after modification by Ch-T. The  $hSlo1 + M7L$ ,  $M23L$ ,  $M177L$ , or Triple  $V_{0.5}$  values for the results obtained before Ch-T application (open circles) were  $153.9 \pm 12.5$  mV ( $n = 4$ ),  $147.1 \pm 3.2$  mV ( $n = 5$ ),  $164.1 \pm 8.3$  mV ( $n = 5$ ), and  $153.8 \pm 6.2$  mV ( $n = 6$ ), respectively. After Ch-T application (closed circles), the  $hSlo1 + M7L$ ,  $M23L$ ,  $M177L$ , or Triple  $V_{0.5}$  values were  $85.5 \pm 9$  mV ( $\Delta V_{0.5}$  range,  $-54$  to  $-78$  mV;  $P = 0.001$ ,  $n = 4$ ),  $96.9 \pm 6.4$  mV ( $\Delta V_{0.5}$  range,  $-42$  to  $-60$  mV;  $P = 0.0002$ ,  $n = 5$ ),  $105 \pm 5.9$  mV ( $\Delta V_{0.5}$  range,  $-43$  to  $-71$  mV;  $P = 0.0005$ ,  $n = 5$ ), and  $103.1 \pm 7.3$  mV ( $\Delta V_{0.5}$  range,  $-45$  to  $-60$  mV;  $P < 0.0001$ ,  $n = 6$ ), respectively. (C) The mean  $\Delta V_{0.5}$  values (left) for  $hSlo1 + M7L$ ,  $M23L$ ,  $M177L$ , and Triple  $m\beta 1$  were  $-68.4 \pm 5.2$  mV,  $-50.2 \pm 4.1$  mV,  $-59.0 \pm 5.8$  mV, and  $-50.7 \pm 2.6$  mV, respectively. The mean  $\Delta Q_{app}$  values (right) for  $hSlo1 + M7L$ ,  $M23L$ ,  $M177L$ , and Triple  $m\beta 1$  were  $-0.11 \pm 0.03e$  ( $P = 0.04$ ,  $n = 4$ ),  $-0.14 \pm 0.08e$  ( $P = 0.15$ ,  $n = 5$ ),  $-0.019 \pm 0.03e$  ( $P = 0.6$ ,  $n = 5$ ), and  $-0.12 \pm 0.04e$  ( $P = 0.02$ ,  $n = 6$ ), respectively. A negative  $\Delta V_{0.5}$  indicates a leftward shift of the G-V curve, and a negative  $\Delta Q_{app}$  value indicates a decrease in the slope of the G-V curve after Ch-T modification.

thesis, we could not readily test its role. M89 is present in  $m\beta 1$  but absent in  $b\beta 1$ . However, both  $m\beta 1$  and  $b\beta 1$  confer to  $hSlo1$  the enhanced Ch-T sensitivity, thereby excluding the critical involvement of M89 (Fig. 4). Thus, M7, M23, and M177 were individually mutated to leucine which is much less susceptible to oxidation by Ch-T than methionine (Ciorba et al., 1997). In addition, a triple  $m\beta 1$  mutant (Fig. 5, Triple) in which M7, M23, and M177 were all replaced by leucine was constructed. When coexpressed with  $hSlo1$ , each  $m\beta 1$  mutant channel complex exhibited currents with wild-type  $\beta 1$ -like characteristics including slower activation and deactivation compared with  $hSlo1$  alone, thus confirming that these  $m\beta 1$  mutants functionally associated with  $hSlo1$ .

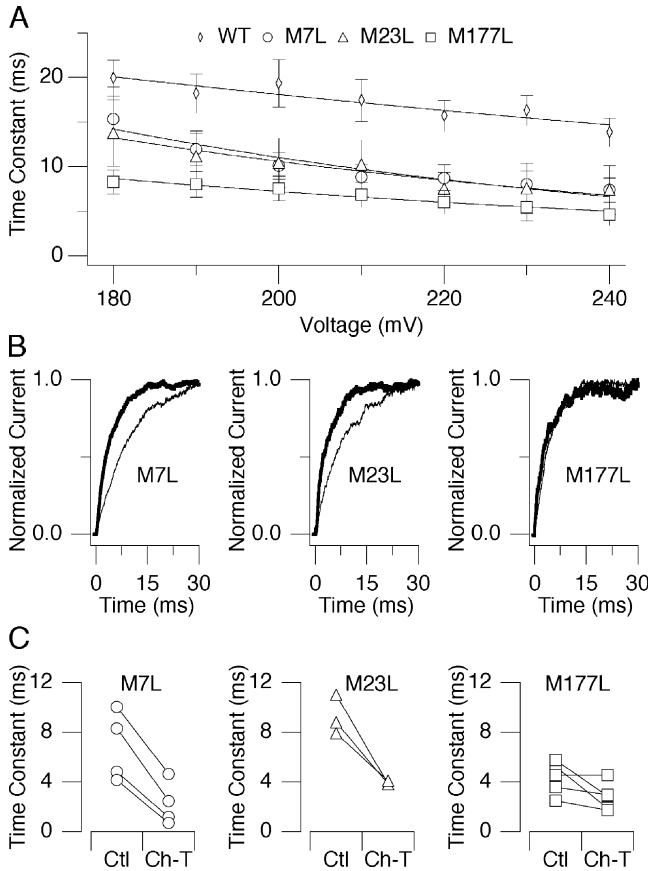
After modification by Ch-T, each  $m\beta 1$  mutant channel complex ( $M7L$ ,  $M23L$ ,  $M177L$ , and Triple) showed a large leftward shift in  $V_{0.5}$  (Fig. 5 B). The mean  $V_{0.5}$  for  $hSlo1 + M7L$ ,  $M23L$ ,  $M177L$ , or Triple  $m\beta 1$  shifted by  $-68 \pm 5.2$  mV,  $-50 \pm 4.1$  mV,  $-59 \pm 5.7$  mV, and  $-50.6 \pm 2.6$  mV, respectively (Fig. 5 C); these  $\Delta V_{0.5}$  were significantly larger than that found with  $hSlo1$  alone ( $\Delta V_{0.5} \approx -30$  mV; Fig. 1 D) ( $P < 0.05$ , Bonferroni test). These results indicated that oxidation of methionine residues within  $m\beta 1$  is not necessary to produce the enhanced G-V curve shift after modification by Ch-T.

#### Methionine Oxidation within $\beta 1$ Is Not Required for the Greater Slowing of $hSlo1$ Deactivation

Treatment with Ch-T slowed the deactivation time course of every  $hSlo1 +$  mutant  $m\beta 1$  complex examined (Fig. 6 A). The extent of this slowing of  $hSlo1$  deactivation



**FIGURE 6.** Slowing of channel deactivation after oxidation by Ch-T is maintained in all  $m\beta 1$  methionine mutant channel complexes. (A) Superimposed  $hSlo1 + M7L$ ,  $M23L$ ,  $M177L$ , or Triple  $m\beta 1$  normalized tail currents recorded at  $-40$  mV after pulses to  $180$  mV before (thin sweeps) and after (thick sweeps) treatment with Ch-T. (B) Voltage dependence of the deactivation time constant for  $hSlo1 +$  wild type (diamonds;  $n = 5$ ),  $M7L$  (circles;  $n = 4$ ),  $M23L$  (triangles;  $n = 5$ ),  $M177L$  (squares;  $n = 5$ ), or Triple (inverted triangles;  $n = 6$ )  $m\beta 1$  before (open symbols) and after (closed symbols) Ch-T treatment.

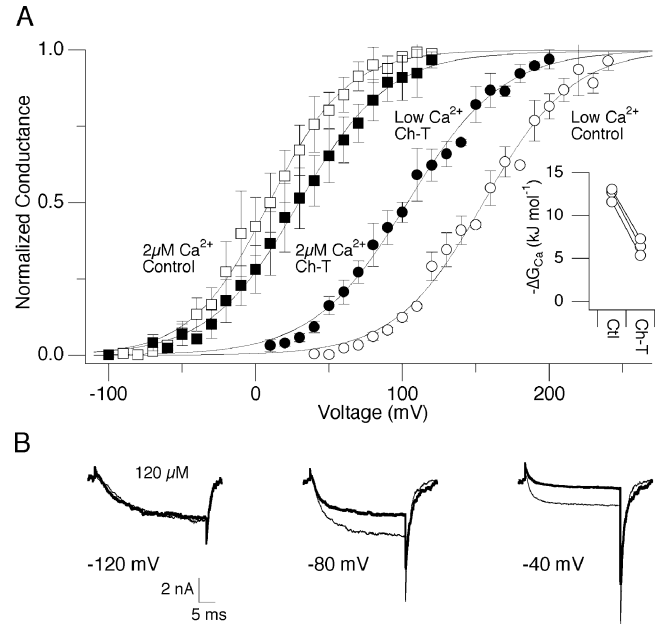


**FIGURE 7.** M177 in  $m\beta 1$  specifically affects channel activation. (A) Voltage dependence of the activation time constant before treatment with Ch-T for  $hSlo1$  + wild type (diamonds;  $n = 14$ ), M7L (circles;  $n = 4$ ), M23L (triangles;  $n = 5$ ), or M177L (squares;  $n = 5$ )  $m\beta 1$ . (B) Currents recorded at 220 mV from the holding voltage of 0 mV before (thin sweep) and after (thick sweep) modification by Ch-T. (C) Activation time constant values at 220 mV before and after oxidation by Ch-T from individual experiments.

tion with any of the  $m\beta 1$  mutants was indistinguishable from the slowing of  $hSlo1$  +  $m\beta 1$  (Fig. 6 B). The voltage dependence of the deactivation kinetics was also unaltered by Ch-T treatment with the equivalent charge movement remaining at  $\sim 0.3e$  in all cases. Thus, the Ch-T-induced slowing of channel deactivation did not specifically require M7, M23, M177, or the presence of all three residues together in  $\beta 1$ .

#### M177 in $m\beta 1$ Is Critical for Typical $hSlo1$ Activation Properties

Oxidative modification by Ch-T accelerated the activation kinetics of  $hSlo1$  +  $m\beta 1$  but not of  $hSlo1$  alone (Fig. 3). For channel complexes that included an  $m\beta 1$  methionine point mutant, there was a trend for the activation kinetics to be faster than  $hSlo1$  + wild-type  $m\beta 1$  even before Ch-T treatment (Fig. 7 A). However, a difference in the activation time course was statistically significant only in  $hSlo1$  + M177L  $m\beta 1$  (220 mV;  $P =$



**FIGURE 8.** The effect of Ch-T on channel open probability depends on  $Ca^{2+}$ . (A) G-V curves before and after modification by Ch-T. Currents were first generated by pulsing to different test potentials from a holding voltage of 0 mV in the virtual absence of  $Ca^{2+}$ . This recording protocol was repeated after bath application of 2.1  $\mu M$   $Ca^{2+}$ . Tail currents were measured at  $-50$  mV in zero  $[Ca^{2+}]_i$  or  $-100$  mV in high  $[Ca^{2+}]_i$ . After a return to zero  $Ca^{2+}$  for treatment with 2 mM Ch-T, the recording protocol was again repeated in zero and then 2.1  $\mu M$   $Ca^{2+}$ . The zero  $[Ca^{2+}]_i$   $V_{0.5}$  values for the results obtained before (open circles) and after (closed circles) Ch-T application were  $155.3 \pm 5.1$  mV and  $103.5 \pm 7.2$  mV ( $\Delta V_{0.5}$  range,  $-47$  to  $-56$  mV;  $P = 0.002$ ,  $n = 3$ ), respectively. The smaller  $\Delta V_{0.5}$  value (approximately  $-50$  vs.  $-75$  mV; Fig. 1) was most likely because of the use of an external recording solution containing less  $K^+$  than that previously used in the zero  $[Ca^{2+}]_i$  experiments. The 2.1  $\mu M$   $[Ca^{2+}]_i$   $V_{0.5}$  values for the results obtained before (open squares) and after (closed squares) Ch-T application were  $9.1 \pm 11.2$  mV and  $29.6 \pm 12.5$  mV ( $\Delta V_{0.5}$  range,  $15$ – $25$  mV;  $P = 0.018$ ,  $n = 3$ ), respectively. The zero  $[Ca^{2+}]_i$   $Q_{app}$  values for the results obtained before and after Ch-T application were  $0.84 \pm 0.04e$  and  $0.79 \pm 0.02e$  ( $P = 0.13$ ,  $n = 3$ ), respectively. The 2.1  $\mu M$   $[Ca^{2+}]_i$   $Q_{app}$  values for the results obtained before and after Ch-T application were  $1.03 \pm 0.07e$  and  $0.88 \pm 0.03e$  ( $P = 0.1$ ,  $n = 3$ ), respectively. (inset) The contribution of  $Ca^{2+}$ -dependent gating to overall channel opening ( $\Delta G_{Ca}$ ) before and after oxidation by Ch-T from individual experiments. (B) Representative currents from a single patch ( $n = 5$ ) in the presence of 120  $\mu M$   $[Ca^{2+}]_i$  before (thin sweeps) and after (thick sweeps) Ch-T treatment. The currents were elicited from a holding voltage of  $-200$  mV.

0.002, Bonferroni test). In fact, the activation kinetics of  $hSlo1$  + M177L  $m\beta 1$  before Ch-T treatment was similar to that of the  $hSlo1$  + wild-type  $m\beta 1$  complex after modification by Ch-T.

The activation kinetics of  $hSlo1$  + M7L  $m\beta 1$  and  $hSlo1$  + M23L  $m\beta 1$  after Ch-T treatment were significantly faster, as found with wild-type  $m\beta 1$  (Fig. 7, B and C;  $P = 0.005$  and  $0.03$ , respectively). However, Ch-T



failed to accelerate the activation time course of *hSlo1* + M177L *mβ1* in an appreciable manner ( $P \geq 0.085$ ). M177L *mβ1* does associate with *hSlo1* because the deactivation kinetics of *hSlo1* + M177L *mβ1* was indistinguishable from *hSlo1* + *mβ1* (Fig. 6 B). Thus, M177 in TM2 of *β1* is a key determinant of the activation kinetics and the oxidative sensitivity of *hSlo1* + *mβ1*.

#### The Effect of Ch-T Treatment with *β1* Present Is $Ca^{2+}$ Dependent

The hyperpolarizing shift in  $V_{0.5}$  caused by treatment with Ch-T was essentially eliminated by increasing  $[Ca^{2+}]_i$  to 2.1  $\mu M$  (Fig. 8 A). Similar results were obtained with saturating levels of divalent ions,  $[Ca^{2+}]_i = 120 \mu M$  and  $[Mg^{2+}]_i = 10 mM$  (Fig. 8 B). In the presence of elevated  $[Ca^{2+}]_i$ , cysteine oxidation is capable of shifting the G-V curve to the right (Tang et al., 2004), which may account for the depolarizing shift seen here after treatment with Ch-T. With the assumption that the free energy changes associated with the  $BK_{Ca}$  channel intrinsic opening process, voltage-dependent activation, and  $Ca^{2+}$  binding together contribute to the overall open probability in a linearly additive manner (Cui and Aldrich, 2000), the measured G-V parameters were used to infer the free energy contributions of  $Ca^{2+}$  to channel opening (Tang et al., 2004) in the control and Ch-T treated conditions. The decrease in  $\Delta G_{Ca}$  ( $\sim 50\%$ ) after Ch-T modification of *hSlo1* + *mβ1* in 2.1  $\mu M$   $[Ca^{2+}]_i$  indicates that  $Ca^{2+}$  makes a smaller free energy contribution to overall channel opening after oxidation (Fig. 8 A, inset).

#### Biophysical Model Simulation

The functional effects of oxidation by Ch-T of the *hSlo1* + *β1* complex in the virtual absence of  $Ca^{2+}$  may be interpreted using the HCA allosteric gating model (Fig. 9 A; Horrigan et al., 1999) as performed for the *hSlo1* channel without *β1* (Tang et al., 2001). The voltage dependence of *hSlo1* alone shifts by  $-30 mV$ , and the deactivation time course slows after oxidation by Ch-T. To account for these alterations, Tang et al. (2001) increased the value of the strongly voltage-dependent parameter  $\alpha(0)$ , which may correspond to movement of the voltage sensor (Horrigan et al., 1999), by 2.3-fold and decreased the rate constant of the closing transition dominant at negative voltages ( $\gamma_0$ ) by 60% (Fig. 9 B; Tang et al., 2001, Fig. 14). We simulated the potentiated effects of Ch-T treatment in the presence of *β1* in the following manner. First, the value of  $\alpha(0)$  is further increased (about twofold) to account for the larger shift,  $-75 mV$ , of the G-V curve (Fig. 9 B). Second, the closing rate constant  $\gamma_0$  decreases by an additional 40% to account for the greater slowing of the tail kinetics. Third, in addition to the two quantitative changes listed, the rate constant for the

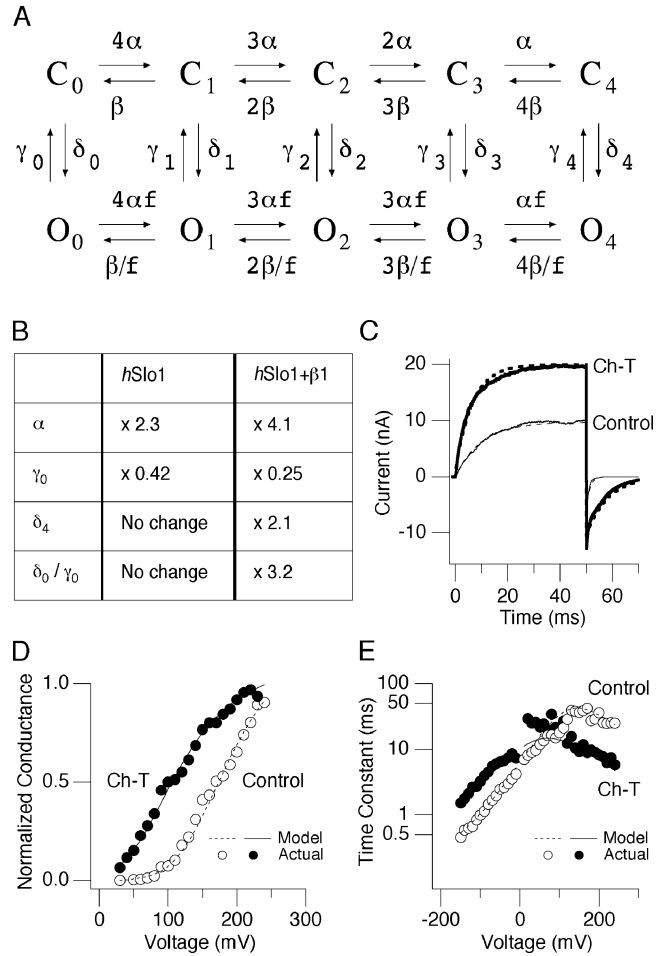


FIGURE 9. Simulation of oxidation by Ch-T on *hSlo1* + *β1* function based on the HCA model. (A) The HCA allosteric gating model (Horrigan et al., 1999). The most probable opening of the channel at strongly depolarized voltages involves transitions from  $C_0$  to  $C_1$ ,  $C_2$ ,  $C_3$ ,  $C_4$ , and then  $O_4$ . Likewise, channel closing at negative voltages entails transitions from  $O_4$  to  $O_3$ ,  $O_2$ ,  $O_1$ ,  $O_0$ , and then  $C_0$ . (B) Adjustments in average parameter values from the HCA model required to simulate the effect of modification by Ch-T on *hSlo1* (Tang et al., 2001) or *hSlo1* + *β1* function. HCA model values are as follows:  $\alpha(0) = 1,500 s^{-1}$ ,  $\beta(0) = 35,370 s^{-1}$ ,  $\delta_0(0) = 0.007 s^{-1}$ ,  $\delta_1(0) = 0.154 s^{-1}$ ,  $\delta_2(0) = 3.39 s^{-1}$ ,  $\delta_3(0) = 52 s^{-1}$ ,  $\delta_4(0) = 65 s^{-1}$ ,  $D = 22$ ,  $f = D^{0.5}$ , and  $L(0) = \delta_0(0)/\gamma_0(0) = 2 \times 10^{-6}$ .  $L(0)$  represents the open-to-closed equilibrium constant in the absence of an applied voltage. (C) Experimental *hSlo1* + *mβ1* currents (continuous sweeps) recorded at 160 mV before (thin) and after (thick) oxidation by Ch-T and simulated currents (dashed lines) from the HCA model adjusted for the effect of Ch-T in the presence of *β1*. (D) G-V curves from a patch containing *hSlo1* + *mβ1* before (open circles) and after (closed circles) treatment with Ch-T. Data simulated from the *hSlo1* + *β1* model before (dotted line) and after (continuous line) Ch-T are superimposed. (E) Activation/deactivation time constants. Symbols are the same as in D.

opening transition dominant at positive voltages ( $\delta_4$ ) is increased by 2.1-fold to account for the unique acceleration of the activation kinetics observed in *hSlo1* + *β1* but not in *hSlo1* alone. Simulated data produced from

this model that account for the effect of Ch-T on *hSlo1* function with  $\beta 1$  present match the experimental data (Fig. 9, C–E).

## DISCUSSION

### *Methionine Oxidation Leads to Distinct Alterations in *hSlo1* + $\beta 1$ Function*

Coexpression of  $\beta 1$  with *hSlo1* is known to affect the activation/deactivation kinetics and apparent  $\text{Ca}^{2+}$  sensitivity of the channel (Orio et al., 2002). Here, we have demonstrated that oxidation of *hSlo1* in the presence of the auxiliary subunit  $\beta 1$  leads to functional effects clearly distinguishable from those observed with *hSlo1* alone. Oxidation of *hSlo1* promoted by Ch-T causes a leftward shift of the G-V curve by  $\sim 30$  mV. However, this hyperpolarizing shift is more than twice as great ( $\sim 75$  mV) in the presence of  $\beta 1$ . Furthermore, the Ch-T-induced slowing of *hSlo1* deactivation is even more dramatic with  $\beta 1$  present. In addition,  $\beta 1$  confers a novel effect of oxidation not observed with *hSlo1* alone; modification by Ch-T leads to the distinct acceleration of *hSlo1* activation evident at each depolarized voltage only with the inclusion of  $\beta 1$  into the channel complex. These unique features of oxidative modification in the presence of the  $\beta 1$  subunit overall cannot be accounted for by a difference in the modification rate as compared with *hSlo1* alone or a simple voltage-dependent shift in the open probability and activation/deactivation kinetics.

### *Role of Cysteine and Methionine Residues in $\beta 1$*

Because Ch-T preferentially oxidizes methionine residues under physiological conditions (Levine et al., 1996), methionine is implicated as the main target of oxidation by Ch-T that is responsible for the observed functional alterations in both *hSlo1* and *hSlo1* +  $\beta 1$ . However, protein-modifying agents may not be perfectly specific for one particular amino acid. In fact, both cysteine and methionine are possible physiological targets of Ch-T. To determine if cysteine oxidation plays a role in the Ch-T effect on *hSlo1* alone, Tang et al. (2001) previously showed that cysteine-specific reagents (5,5'-dithio-bis (2-nitrobenzoic acid), methanethiosulfonate ethylammonium, and *p*-chloromercuribenzoic acid) actually decreased channel activity, thereby demonstrating that cysteine oxidation has opposite effects on channel function than methionine oxidation. Furthermore, the Ch-T-induced potentiation was maintained in *hSlo1* mutants that lacked most of the cysteine residues within the channel. Finally, peptide methionine sulfoxide reductase, an enzyme that catalyzes the reduction of met-O (Weissbach et al., 2002), partially reversed the effect of Ch-T treatment. Therefore, the functional alterations caused by

Ch-T were attributed to methionine oxidation within *hSlo1*.

Cysteine residues within  $\beta 1$  are not required for typical regulation of *hSlo1* kinetics or the enhanced effects on channel function after oxidation. The *b* $\beta 1$  subunit devoid of any cysteine residues behaves much like wild-type *b* $\beta 1$  in terms of slowing *hSlo1* activation and deactivation. Furthermore, after oxidation by Ch-T, the cysteine mutations still permit the significantly larger  $\Delta V_{0.5}$  value, the slower deactivation kinetics and the accelerated channel activation similar to those observed with wild-type *b* $\beta 1$ . These results indicate that cysteine is not the likely Ch-T target responsible for causing the functional changes after oxidation.

Similar to the *b* $\beta 1$  cysteine mutant, the *m* $\beta 1$  methionine mutants regulate *hSlo1* kinetics much like wild-type  $\beta 1$ . Moreover, all *m* $\beta 1$  methionine mutants including the triple mutant maintain the dramatic shift of the G-V curve toward the hyperpolarizing direction and the enhanced slowing of channel deactivation after oxidation by Ch-T. Evaluation of the role of the initial  $\beta 1$  methionine residue (M1) is not straightforward. However, its contribution to the enhanced oxidative regulation of *hSlo1*, although a possibility, is unlikely due to potential removal by posttranslational processing of the mature protein (Creighton, 1993). Therefore, the presence of the  $\beta 1$  subunit provides the possibility to amplify the functional effects of methionine oxidation within the *hSlo1* pore-forming subunit with regard to channel open probability and deactivation.

### *$\beta 1$ M177 Involvement in the Functional Interaction with *Slo1**

Although the enhanced shift of the G-V curve and slowing of *hSlo1* deactivation does not require cysteine or methionine residues within  $\beta 1$ , the effect of oxidation on *hSlo1* activation critically depends on M177 in TM2 of *m* $\beta 1$ . In the control condition, only M177L *m* $\beta 1$  causes a significant difference in the channel activation time course. Furthermore, the M177L mutation eliminates the oxidative sensitivity of channel activation typically observed with  $\beta 1$  present. Thus, M177 controls the *hSlo1* activation kinetics at very positive voltages, which is described by the rate constant  $\delta_4$  in the HCA model (Fig. 9), and oxidation of M177 to met-O most likely mediates the Ch-T-induced acceleration of activation kinetics. However, the possibility that the M177L mutation hinders the access of Ch-T to its target elsewhere cannot be completely eliminated.

The mutant-specific effect on channel activation suggests a partial uncoupling of *hSlo1* and  $\beta 1$  because of mutation or oxidation at the M177 position. Because the activation kinetics of *hSlo1* is faster without  $\beta 1$ , oxidation of *hSlo1* +  $\beta 1$  may cause channel activation to be more like *hSlo1* alone by removal of the  $\beta 1$  influ-

ence. The hydrophobic leucine mutation at M177 mimics the presence of met-O at that location because the control activation kinetics of *hSlo1* + M177L *mβ1* resembles that of oxidized *hSlo1* + wild-type *mβ1*. Indeed, an increase in surface hydrophobicity, while somewhat paradoxical, has been shown after oxidation of methionine residues within the enzyme glutamine synthetase (Levine et al., 1996). Perhaps oxidation of M177 to met-O, whereby acting as the sensor or switch, partially disrupts an interaction between the  $\beta 1$  subunit and the structural elements in or near the RCK (regulator of  $K^+$  conductance) domains within *hSlo1* that are specifically responsible for controlling activation kinetics. Similar to the effect of  $\beta 1$  M177 on channel activation, other residues within different  $\beta$  subunits influence functional coupling of the auxiliary subunit and *hSlo1*. For example, the phosphorylation states of T11/S17 in the cytoplasmic  $NH_2$  terminus and S210 in the cytoplasmic COOH terminus within  $\beta 4$  affect the functional coupling between *hSlo1* and  $\beta 4$ , as determined by changes in channel voltage dependence and activation/deactivation kinetics specific to modification of the different residues (Jin et al., 2002).

#### *Physiological Implications*

As found with *hSlo1* alone, the effect of methionine oxidation on *hSlo1* function in the presence of  $\beta 1$  is sensitive to  $[Ca^{2+}]_i$ . In the virtual absence of  $[Ca^{2+}]_i$ , *hSlo1* + *mβ1* displays a hyperpolarizing shift of  $V_{0.5}$  that is twice as great as *hSlo1* alone after oxidation by Ch-T. This Ch-T-induced shift resembles the presence of  $\sim 0.4 \mu M$   $[Ca^{2+}]_i$  (Cox and Aldrich, 2000). The oxidized channel complex can open in the physiological voltage range ( $< 50$  mV) without  $[Ca^{2+}]_i$  as further evidenced by the increase in open probability observed at  $-40$  mV. An increase in channel open probability at low  $[Ca^{2+}]_i$  could have an impact on resting  $BK_{Ca}$  channel activity in smooth muscle cells, thereby influencing vascular tone. Because  $BK_{Ca}$  channels crucially shape the action potential posthyperpolarization phase in certain cell types, this increase in channel open probability may prevent unregulated neuronal firing (Lancaster and Nicoll, 1987; Storm, 1987; Marsh and Brown, 1991; Zhang and McBain, 1995; Pedarzani et al., 2000; Faber and Sah, 2002; Edgerton and Reinhart, 2003).

The concept that the binding of  $Ca^{2+}$  performs mechanical work to open the Slo1 channel (Jiang et al., 2002) has been further developed into a spring-based gating mechanism in which the diameter of the gating ring, formed by the RCK domains from each Slo1 subunit, expands upon  $Ca^{2+}$  binding, thereby generating an active force that pulls the S6-RCK1 linker regions that act as the springs, thus opening the channel gates (Niu et al., 2004). This proposed gating process might be similarly affected by methionine oxidation, which bi-

ases the open channel state. In the absence of  $[Ca^{2+}]_i$ , oxidation of methionine residues to met-O within the *hSlo1* pore-forming subunit may likewise affect the structure or position of the gating ring ultimately influencing gating of the channel. The lack of a hyperpolarizing shift of  $V_{0.5}$  in response to modification by Ch-T at high  $[Ca^{2+}]_i$  indicates that the effects of  $Ca^{2+}$  and methionine oxidation on channel gating are not additive and may in fact operate on the same effectors. In the case of the *hSlo1* +  $\beta 1$  channel complex, the presence of  $\beta 1$  may cause a unique conformational change in *hSlo1*, such that additional methionine residues in *hSlo1* are exposed and able to react with Ch-T, thereby accounting for the enhanced functional effects of oxidation. However, the similarity in the modification time courses of *hSlo1* and *hSlo1* + *mβ1* argues against this possibility.

Modification of ion channels by ROS/RNS during oxidative stress could alter channel function and eventually disrupt normal  $[Ca^{2+}]_i$  and other homeostatic parameters (Kourie, 1998). Potential consequences of oxidative stress include accelerated aging (Hensley and Floyd, 2002), as well as pathophysiological conditions such as various neurodegenerative disorders (Coyle and Puttfarcken, 1993) and ischemia-reperfusion injury after stroke (Babbs, 1988; Rubanyi, 1988). However, certain ion channel modifications by ROS/RNS may serve as compensatory mechanisms to oxidative assault. One such example involves the mitochondrial ATP-sensitive  $K^+$  channel ( $mitoK_{ATP}$ ) that is activated by ROS during initial, mild ischemia; as a result, the heart is preconditioned to future ischemic attacks and infarctions (Szewczyk and Marban, 1999; Grover and Garlid, 2000; Zhang et al., 2001). Much like the  $mitoK_{ATP}$  channel, the  $BK_{Ca}$  channel clearly represents a prime candidate for aiding in the recovery from ROS/RNS attack given its localization in brain and smooth muscle, as well as the documented oxidation-related alteration of its function (DiChiara and Reinhart, 1997; Sobey et al., 1997; Wang and Wu, 1997; Wang et al., 1997; Barlow et al., 2000; Gong et al., 2000; Soh et al., 2001; Tang et al., 2001, 2004; Brakemeier et al., 2003). Whether the  $BK_{Ca}$  channel contributes to the progression of oxidative stress-related conditions or instead serves a more compensatory role—such as maintaining resting membrane potential if  $[Ca^{2+}]_i$  is disrupted—remains to be determined.

In summary, we showed that in the virtual absence of  $Ca^{2+}$ , methionine oxidation by Ch-T dramatically alters *hSlo1* function with the association of the  $\beta 1$  subunit. The presence of  $\beta 1$  as opposed to methionine and/or cysteine oxidation within this auxiliary subunit greatly amplifies the increase in channel open probability and the slowing of deactivation derived from oxidation of the *hSlo1* pore-forming subunit. The target methionine

residues within *hSlo1* are not yet known, but may be found in the S5/P/S6 segments (Tang et al., 2001) and/or the gating ring region (Niu et al., 2004). In contrast, M177 within  $\beta 1$  influences *hSlo1* activation and most likely serves as the methionine target responsible for the acceleration in channel activation after methionine oxidation in the presence of the  $\beta 1$  subunit. Testing the oxidative effects with  $\beta$  subunits present provides more relevant results that can then be readily extended to physiological or pathophysiological conditions. Whether the effect of methionine oxidation on *hSlo* function in the presence of other  $\beta$  subunits ( $\beta 2$ – $4$ ) also occurs remains to be determined, but  $\beta 1$  clearly facilitates unique modulation of channel function in the face of oxidation.

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