Outward Currents Influencing Bursting Dynamics in Guinea Pig Trigeminal Motoneurons

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INTRODUCTION

Neuronal bursting behavior, by a traditional mechanism (Del Negro et al. 1998), utilizes non- or slowly inactivating inward currents to generate a region of negative slope resistance in the steady-state current-voltage (I-V) relationship. This region of negative slope resistance creates two stable membrane states on either side of spike threshold: one quiescent state and another superthreshold or active state, at which the cell discharges tonically. These two states are traversed during the quiescent and active phases of bursting.

To initiate individual bursts, the inward currents underlying the negative slope resistance cause rapid depolarization, which shifts the membrane potential from its quiescent to active state (see Fig. 3B of Del Negro et al. 1998). Conversely, bursts terminate because of much slower membrane processes that generate repolarizing currents (Bertram et al. 1995; Rinzel and Ermentrout 1998). During the active phase these steadily growing currents cause a decline in intraburst spike frequency and ultimately a return to quiescence. The source of the repolarizing currents can be a distinct set of ion channels, intramembranous ion pumps, the slow inactivation of inward currents, or a combination of these factors. The repolarizing currents that terminate bursts are usually activated by cytosolic cation accumulation (Gorman et al. 1981, 1982; Li and Hatton 1996; Rose and Ransom 1997). For example, inward Ca2+ current can activate Ca2+-dependent K+ currents during bursts (Lengendre et al. 1985; Tell and Jean 1993) or cause Ca2+-dependent Ca2+ current inactivation (Benson and Adams 1989; Canavier et al. 1991; Heyer and Lux 1976; Kramer and Zucker 1985). Na+ currents can activate the Na+/K+-ATPase pump during bursts (Johnson et al. 1992; Li et al. 1996; Tsai and Chen 1995) or possibly experience voltage- or cytosolic Na+-dependent inactivation (Fleidervish et al. 1996).

In trigeminal motoneurons that innervate jaw musculature both Ca2+ and Na+ currents are important for serotonin-induced bursting (Hsiao et al. 1998). Therefore intracellular accumulation of Ca2+ and Na+ ions may activate distinct slow repolarizing currents. Here we test the hypothesis that an apamin-sensitive, Ca2+-dependent K+ current (IK,Ca) and a cardiac glycoside-sensitive Na+/K+-ATPase pump current (INa) activated by intracellular Ca2+ and Na+ ions, respectively, contribute to burst termination and the overall dynamics of bursting behavior.

METHODS

Trigeminal motoneurons were recorded intracellularly in 450-μm-thick transverse brain stem slices from adolescent guinea pigs (150–250 g), as shown previously (Chandler et al. 1994; Hsiao et al. 1998). Slices were perfused in a gas-interface chamber at 34°C with oxygenated Ringer solution containing (in mM) 130.0 NaCl, 3.0 KCl, 1.25 KH2PO4, 20.0 NaHCO3, 10.0 d-glucose, 2.4 CaCl2, and 1.3 MgSO4. Microelectrodes were filled with 3 M KCl (15–20 MΩ). Current-clamp experiments were performed with an Axoclamp 2A amplifier in bridge mode (Axon Instruments, Burlingame, CA).

The acceptance criteria for trigeminal motoneurons included resting potential less than ~50 mV, input resistance of 6–18 MΩ, and action potential amplitude >60 mV (Chandler et al. 1994; Curtis and Appenteng 1993).

Bath-applied drugs were effective after ~3.5 min of perfusion; complete solution change required ~20 min. We used 10 μM serotonin (Sigma Chemical, St. Louis, MO) to elicit bursting; this dose reliably induces bistability and bursting in trigeminal motoneurons (Hsiao et al. 1998). Other drugs were used at the following concentrations: apamin (0.2 μM, Sigma), ouabain (6 μM, Sigma), and strophanthidin (4 μM, Sigma). In some experiments Ca2+ concentration was lowered to 0.4 mM and substituted with Ba2+ on an equimolar basis.

Bursting characteristics such as the time of burst onset and termination and intraburst spike frequency were measured with Datapac III v0.1.47 (Run Technologies, Irvine, CA). Burst duration was defined as the mean time from burst onset to termination in three or more consecutive bursts. Burst cycle time was defined as the mean time from burst onset to onset in three or more consecutive cycles. Means...
mV compared with burst onset, suggesting the growth of termination, the membrane potential was hyperpolarized by 5 repolarizing current (Fig. 1, failed to reach threshold putatively because of the growing that the voltage trajectory rose after the previous spike but izing afterpotential occurred at burst termination. This suggests et al. 1995; Del Negro et al. 1998). Also a hump-like depolar- negative slope resistance-based bursting mechanism (Bertram phase. First, during the plateau potentials and bursts, spike repolarizing currents were recruited throughout the active cycle time ranged from 4.9 to 178 s.

5 s, the interburst interval ranged from 4.2 to 123 s, and the 90). In all cells recorded, burst duration ranged from 0.7 to 202 mV), plateau potentials could be evoked by 300-ms, 1-nA current pulses (bottom trace). Continuous bursting activity was evoked by adding a constant current bias of 0.5 nA. By removing the depolarizing current bias, the cell reestablished its resting membrane potential and again exhibited plateau potentials in response to the transient stimuli. The current calibration applies to A and B. The time calibration applies only to A, B, and C; plateau potential (B) and a burst (C) from the record in A (responses labeled 1 and 2, respectively) are displayed with an expanded time scale. For each response, the instantaneous intraburst spike frequency is plotted synchronously below the voltage trajectory. The voltage calibration in C applies to all traces. Time calibrations for B and C are displayed as the abscissa of the frequency-time (f-t) plot (bottom).

are reported as ±SE. Pharmacologically induced changes in bursting characteristics were tested for each cell individually with unpaired Student’s t-tests (SYSTAT 7.0, SPSS, Chicago, IL).

R E S U L T S

In the presence of serotonin, trigeminal motoneurons exhibit plateau potentials and spontaneous rhythmic bursting (Hsiao et al. 1998). Figure 1 illustrates these behaviors in a typical cell. At resting potential, the cell was bistable. Therefore transient stimuli evoked plateau potentials (Fig. 1, A and B, n = 135, including cells from our earlier paper) (Hsiao et al. 1998). This behavior could be converted to continuous bursting activity by injecting depolarizing constant current (Fig. 1, A and C, n = 90). In all cells recorded, burst duration ranged from 0.7 to 53 s, the interburst interval ranged from 4.2 to 123 s, and the cycle time ranged from 4.9 to 178 s.

Several features of the plateau potentials and bursts suggest that repolarizing currents were recruited throughout the active phase. First, during the plateau potentials and bursts, spike frequency declined monotonically until burst termination (Fig. 1, B and C, bottom), which is characteristic of a traditional, negative slope resistance-based bursting mechanism (Bertram et al. 1995; Del Negro et al. 1998). Also a hump-like depolarizing afterpotential occurred at burst termination. This suggests that the voltage trajectory rose after the previous spike but failed to reach threshold putatively because of the growing repolarizing current (Fig. 1, B and C, arrows). Last, after burst termination, the membrane potential was hyperpolarized by 5 mV compared with burst onset, suggesting the growth of outward currents during the burst (Fig. 1C, broken line inserted for reference).

Role of Ca$^{2+}$-dependent K$^+$ current

L-type Ca$^{2+}$ currents provide the majority of inward current needed for serotonin-induced bursting (Hsiao et al. 1998). Because Ca$^{2+}$ may accumulate intracellularly during bursts and trigeminal motoneurons express an apamin-sensitive, Ca$^{2+}$-dependent K$^+$ current (Chandler et al. 1994; Kobayashi et al. 1997), we tested whether $I_{K_{Ca}}$ was involved in the burst termination. Specifically, we applied apamin (n = 5) or Ba$^{2+}$-substituted, low-Ca$^{2+}$ solution (n = 4) to reduce or eliminate $I_{K_{Ca}}$.

Apamin typically prolonged burst duration and extended the cycle time. In Fig. 2A mean burst duration increased significantly from 47 ± 2 to 271 ± 30 s (P < 0.001), and mean cycle time increased from 115 ± 5 to 490 ± 46 s (P < 0.001) in the presence of apamin. In general, apamin significantly increased burst duration by 313 ± 76% (n = 5/5, P < 0.01) and extended burst cycle time by 202 ± 58% (n = 4/5, P < 0.05). In addition to increasing burst duration and cycle time, apamin...
also elevated the intraburst spike frequency-time (f-t) relationship (Fig. 2C), consistent with its suppression of the medium-duration afterhyperpolarization (mAHP) in trigeminal motoneurons (Fig. 3B) (Chandler et al. 1994; Hsiao et al. 1997; Kobayashi et al. 1997).

We also tested the role of $I_{KCa}$ with Ba$^{2+}$-substituted, low-Ca$^{2+}$ solution because Ba$^{2+}$ can penetrate Ca$^{2+}$ channels but does not activate $I_{KCa}$. Similar to apamin, low-Ca$^{2+}$/Ba$^{2+}$ solution enhanced burst duration and cycle time. In Fig. 2B mean burst duration increased significantly from 3.5 ± 0.2 to 11.9 ± 0.6 s ($P < 0.001$), and mean cycle time increased from 9.0 ± 0.5 to 22.0 ± 0.8 s ($P < 0.001$). In general, low Ca$^{2+}$/Ba$^{2+}$ significantly increased burst duration by 357 ± 142% ($n = 3/4$, $P < 0.01$) and extended cycle time by 271 ± 76% ($n = 3/4$, $P < 0.001$). Like apamin, low-Ca$^{2+}$/Ba$^{2+}$ solution also elevated the intraburst f-t curve (Fig. 2D). The elevation in spike frequency could be caused by suppression of the mAHP, blockade of Ba$^{2+}$-sensitive K$^+$ channels (Hsiao et al. 1997), or a combination of both effects.

Although apamin and low-Ca$^{2+}$/Ba$^{2+}$ solution upwardly shifted the intraburst f-t relationship, bursts still terminated (in the absence of $I_{KCa}$) when the instantaneous intraburst spike frequency declined to low values similar to control. In Fig. 2C, spike frequency at the time of burst termination was 5 Hz in control and 8 Hz in the presence of apamin. In Fig. 2D, the spike frequency was 3 Hz in control and 5 Hz in low-Ca$^{2+}$/Ba$^{2+}$ solution. These data suggest that the net outward current at the time of burst termination was similar in the presence or absence of $I_{KCa}$. Therefore some compensatory mechanism(s) must be present to terminate bursts after the blockade of $I_{KCa}$.

**Role of Na$^+/K^+$-ATPase pump**

If $I_{KCa}$ was the only outward current controlling spike frequency decline and burst termination, we would expect apamin or low-Ca$^{2+}$/Ba$^{2+}$ solution to induce tonic spiking (because bursts would not be able to terminate). However, bursting continued in apamin or low-Ca$^{2+}$/Ba$^{2+}$ conditions, suggesting additional sources of slow repolarizing current. Because transient and persistent Na$^+$ currents contribute to serotonin-induced bursting in trigeminal motoneurons (Hsiao et al. 1998), cytosolic Na$^+$ accumulation might also activate slow repolarizing current, analogous to Ca$^{2+}$, or cause slow Na$^+$ current inactivation. The Na$^+/K^+$-ATPase pump is associated with bursting behavior (Johnson et al. 1992; Li et al. 1996; Tsai and Chen 1995), is activated by cytosolic Na$^+$, and is electrogenic (because of pump stoichiometry: 3 Na$^+$ ions extruded for every 2 K$^+$ ions that enter) (Läuger 1991). Therefore we tested the potential contribution of the Na$^+/K^+$-ATPase pump current ($I_p$) with cardiac glycocides ouabain ($n = 8$) and strophanthinid ($n = 6$), which block pump activity.

To determine the role of $I_p$ we first applied apamin ($n = 4$, Fig. 3A) or low-Ca$^{2+}$/Ba$^{2+}$ solution ($n = 4$, Fig. 4A) to block $I_{KCa}$. Once apamin or low Ca$^{2+}$/Ba$^{2+}$ was effective, as measured by blockade of the mAhP (Fig. 3B, top), ouabain was added to block $I_p$. In all trigeminal motoneurons tested, ouabain ultimately abolished bursting and induced tonic spiking. Nevertheless, in three of four cells exposed to apamin + ouabain (Fig. 3C) and two of four cells exposed to low Ca$^{2+}$/Ba$^{2+}$ + ouabain (Fig. 4A), bursting activity continued during a transition period that preceded tonic spiking.

During this transitional bursting phase, ouabain decreased burst duration and cycle time and reduced the amplitude of the postburst afterhyperpolarization (AHP). In Fig. 3, burst duration decreased significantly from 10.1 ± 1.0 to 7.6 ± 0.4 s ($P < 0.05$), and cycle time decreased from 29.8 ± 5.2 to 10.6 ± 0.3 s ($P < 0.001$) (measured at the temporal midpoint between the time of ouabain application and the time at which motoneurons assumed tonic spiking). Also ouabain reduced the postburst AHP, as measured during the interburst interval (Fig. 3C, broken line inserted for reference). In low-Ca$^{2+}$/Ba$^{2+}$ solution, ouabain similarly decreased burst duration and cycle time and reduced the postburst AHP (Fig. 4A). In general, ouabain significantly decreased burst duration to 54 ± 11% of burst duration in the presence of apamin alone ($n = 3/3$, $P < 0.05$) and to 68% of burst duration in low-Ca$^{2+}$/Ba$^{2+}$ solution ($n = 1/2$, $P < 0.05$). Moreover, ouabain decreased cycle time to 6 ± 2% of cycle time in the presence of apamin alone ($n = 3/3$, $P < 0.001$) and to 33% of cycle time in low-Ca$^{2+}$/Ba$^{2+}$ solution ($n = 2/2$, $P < 0.001$).

Ouabain ultimately eliminated bursting. When $I_{KCa}$ alone was blocked the intraburst f-t curve shifted upward but re-
remained smooth and monotonic (Figs. 2, C and D, and 3A). When ouabain was added burst duration, burst cycle time, and the postburst AHP all decreased. Later in the experiment ouabain disrupted the smooth monotonic decline in intraburst spike frequency, the hallmark of stable bursting activity. The intraburst f-t curve became erratic and then broke down completely (Fig. 3, Cb–d). Finally, the combined block of \( I_{K,CS} \) and \( I_p \) prevented the rhythmic alternation of active and quiescent bursting phases. Instead, trigeminal motoneurons assumed the tonic spiking state shown in Figs. 3Cd and 4A (middle panel). This suggests that cells entered an active phase that could not terminate, putatively caused by blockade of both slow repolarizing currents, \( I_{K,CS} \) and \( I_p \).

During the nonterminating active phase, action potentials declined slightly in amplitude, \(-20\%\) (Fig. 3B, bottom), and repolarized more slowly, suggesting that ouabain affected the \( \text{Na}^+ \) and \( \text{K}^+ \) concentration gradients to some extent. In the remaining cells exposed to apamin + ouabain (\( n = 1/4 \)) and low-\( \text{Ca}^{2+}/\text{Ba}^{2+} \) + ouabain (\( n = 2/4 \)) the transition to tonic spiking occurred rapidly, and a transient period of bursting activity was indecipherable (not shown).

To further test the role of \( I_p \), we repeated these experiments with the cardiac glycoside strophanthidin. This drug was applied sequentially, after apamin, with the same criteria as the ouabain experiments (\( n = 5/4 \)). Strophanthidin, like ouabain, ultimately abolished bursting and induced a nonterminating active phase (Fig. 4B). Three of six cells exposed to strophanthidin exhibited an analyzable transitional bursting phase. Similar to ouabain, strophanthidin decreased burst duration and cycle time. In Fig. 4B, burst duration decreased significantly from 4.4 \( \pm \) 0.2 to 1.5 \( \pm \) 0.2 s (\( P < 0.01 \)), and cycle time decreased from 11.3 \( \pm \) 0.8 to 2.9 \( \pm \) 0.2 s (\( P < 0.001 \)) (measured at the temporal midpoint between the time of strophanthidin application and the time at which cells assumed tonic spiking). Strophanthidin, similar to ouabain, reduced the postburst AHP (Fig. 4B, broken line inserted for reference). In general, strophanthidin significantly decreased burst duration to 54\% of burst duration in the presence of apamin alone (\( n = 2/3, P < 0.001 \)) and decreased cycle time to 43\% of cycle time in apamin alone (\( n = 3, P < 0.001 \)). One-half of the trigeminal motoneurons exposed to strophanthidin (\( n = 3/6 \)) rapidly assumed tonic spiking without a discernable transitional bursting phase (not shown).

The nonterminating active phase induced by blockade of \( I_{K,CS} \) and \( I_p \) could be manually terminated by extrinsic current injection (\( n = 8 \)). Figure 5 shows a typical cell in the presence of apamin + strophanthidin. Under these conditions rapid burst termination only occurred after repolarizing current injection, whereas bursts initiated spontaneously 10–20 s after readjustment of current bias. These data suggest that the motoneurons still possessed a region of negative slope resistance in the steady-state I-V curve and could initiate bursts (the relationship between the negative slope resistance and burst initiation is explained by Del Negro et al. 1998) but did not possess sufficient intrinsic slow repolarizing processes to terminate bursts autonomously.

To investigate how \( I_p \) influences intraburst spike frequency and the membrane potential trajectory after burst termination, we performed the following experiment. In the absence of serotonin, we applied apamin to trigeminal motoneurons and simulated bursts by applying current pulses of variable magnitude (2–4 nA) and duration (0.4–13.6 s). For each simulated burst, we examined 1) the postburst AHP after termination of the stimulus pulse and 2) the intraburst f-t relationship. We then compared these measures in control versus ouabain (\( n = 6 \)) or strophanthidin (\( n = 1 \)) conditions. In all cells tested ouabain blocked the postburst AHP (Fig. 6A, inset), suggesting that \( I_p \) activated during the simulated burst and produced the ouabain-sensitive AHP at burst termination. Ouabain also shifted the intraburst f-t relationship upward, which further increased the amplitude and latency of the postburst AHP.
bistable or tonic spiking, separated in the \(I-V\) relationship so that the region of negative slope resistance lies below 1.25 mV. Voltage, current, and time calibrations are shown. \(I\): postburst afterhyperpolarization for control and ouabain conditions (superimposed from top traces) in an expanded scale; separate voltage and time calibrations are shown. B: \(f-t\) plots comparing intraburst spike frequency in control (○) and ouabain (●) conditions. Insert: same data on an expanded ordinate axis. Serotonin was not employed for this experiment.

suggests that \(I_p\) normally activated during the simulated burst and constrained intraburst spike frequency (Fig. 6B).

**Discussion**

We examined slow outward currents activated by cytosolic cation accumulation that terminate serotonin-induced bursts in trigeminal motoneurons. We found that \(I_{KCa}\) and \(I_p\) both substantially contribute to burst termination and influence temporal bursting characteristics such as burst duration and the interburst interval. Therefore \(I_{KCa}\) and \(I_p\) probably influence the motoneuronal burst-like patterns that occur during rhythmic oral-motor behaviors (Goldberg and Chandler 1990; Goldberg et al. 1982; Gurahian et al. 1989).

During bursting by a traditional mechanism (Del Negro et al. 1998) neurons phasically alternate between quiescent and active membrane states. Stable membrane states are formed whenever the sum of intrinsic inward and outward currents equals zero and slope conductance is positive. This occurs where the steady-state \(I-V\) relationship intersects its zero-current axis with positive slope. If such an intersection occurs at a voltage less than spike threshold that state is quiescent; if it occurs at a voltage greater than threshold tonic spiking results. To simultaneously possess two stable states neurons must express a region of negative slope resistance in their steady-state \(I-V\) relationship, which can twice intersect the zero-current axis with positive slope.

The inward currents underlying the region of negative slope resistance provide the structural framework for bistability (plateau potentials) and bursting by creating stable membrane states on either side of spike threshold, quiescent and tonic spiking, separated in the \(I-V\) relationship by the region of regenerative inward current. Therefore how do bistable or bursting cells move from state to state; what controls the dynamics?

To answer this, we address 1) how membrane potential first moves from its quiescent to active state, thereby initiating a plateau potential or burst, and 2) what slow repolarizing processes activate during the burst to diminish intraburst spike frequency and ultimately terminate the active phase.

**Burst initiation**

Trigeminal motoneurons utilize L-type \(Ca^{2+}\) and persistent \(Na^+\) currents to initiate and maintain plateau potentials and bursts (Hsiao et al. 1998). Serotonin transforms membrane properties by inducing the necessary region of negative slope resistance and negatively shifting the \(I-V\) relationship so that the region of negative slope resistance often lies below the zero-current axis in the region of net inward current (Chandler and Trueblood 1995; Hsiao et al. 1998). In this case the only stable membrane state is the tonic spiking state, at a depolarized membrane potential. Therefore bursts initiate autonomously (Figs. 1C and 2 of Del Negro et al. 1998).

Sometimes the effects of serotonin are insufficient to shift the region of negative slope resistance fully into the region of net inward current. In this case, the \(I-V\) curve intersects the zero-current axis three times (twice with positive slope), and the cell is bistable. Then from the resting state plateau potentials can be initiated by transient stimuli (e.g., Fig. 1B). Adding depolarizing current bias can extrinsically shift the \(I-V\) relationship so that the negative slope region lies in the area of net inward current, transforming bistability into bursting, as shown in Fig. 1A.

**Burst termination: the role of \(I_{KCa}\)**

Of the slow repolarizing processes that terminate plateau potentials and bursts in trigeminal motoneurons, we first identified \(I_{KCa}\). Because the blockade of \(I_{KCa}\) (by using apamin or low-\(Ca^{2+}/Ba^{2+}\) solution) lengthened burst duration and upwardly shifted the intraburst \(f-t\) curve, we propose that \(I_{KCa}\) normally activates during bursts caused by \(Ca^{2+}\) influx and accumulation, constrains intraburst spike frequency, and helps expedite burst termination. These effects are consistent with how \(I_{KCa}\) is recruited and affects spike frequency in a variety of mammalian motoneurons (Hounsgaard et al. 1988b; Mosfeldt and Rekling 1989; Nishimura et al. 1989; Takahashi 1990; Viana et al. 1993; Zhang and Krmjevic 1987), including trigeminal motoneurons (Chandler et al. 1994; Kobayashi et al. 1997). \(I_{KCa}\) similarly contributes to burst termination during conditional bursting in several other cell types (el Manira et al. 1994; Hu and Bourque 1992; Legendre et al. 1985; Tell and Jean 1993; Wallen and Grillner 1987), including \(N\)-methyl-d-aspartate (NMDA)-induced bursting in trigeminal motoneurons (Kim and Chandler 1995).
However, additional slow processes must influence burst termination in trigeminal motoneurons. When $I_{\text{K,Ca}}$ was blocked bursting activity continued. In the absence of $I_{\text{K,Ca}}$, spike frequency was 5–8 Hz at the time of burst termination, which was similar to the spike frequency at the time of burst termination in control (3–5 Hz, Fig. 2). This suggests that in all conditions the net outward current was comparable at the end of the active phase. Although $I_{\text{K,Ca}}$ normally contributes to burst termination, additional repolarizing processes compensate for $I_{\text{K,Ca}}$ loss under apamin or low-Ca$^{2+}$/Ba$^{2+}$ conditions and generate sufficient repolarizing current to terminate the active phase. However, when $I_{\text{K,Ca}}$ is reduced or eliminated the remaining slow processes require more time to generate the requisite outward current, consequently lengthening burst duration. The extended cycle time during blockade of $I_{\text{K,Ca}}$ most likely reflects the slower time course of relaxation of these compensatory slow repolarizing processes.

**Burst termination: the role of $I_p$**

The other slow repolarizing process, which assists $I_{\text{K,Ca}}$ to terminate plateau potentials and bursts, is constituted primarily by $I_p$ because removal of $I_p$ ultimately blocks serotonin-induced bursting. If slow inactivation of either the persistent Na$^+$ or the L-type Ca$^{2+}$ channels contributed substantially to burst termination we would expect bursts to terminate even when $I_{\text{K,Ca}}$ and $I_p$ are both blocked. However, the combined block of $I_{\text{K,Ca}}$ and $I_p$ always transformed rhythmic bursting into a nonterminating active phase of tonic spiking (Figs. 3 and 4).

We propose that the role of $I_p$ is multifaceted and that the pump current is composed of both steady-state and dynamic components. We base our conclusions on the effects of the cardiac glycosides during the transitional bursting phase that preceded tonic spiking in some cells. First, cardiac glycosides dramatically decreased burst cycle time. Neurons constantly extrude Na$^+$ via the Na$^+$/K$^+$ pump (Läuger 1991). Therefore the electrogenic nature of the pump normally creates a “background” outward current. This steady-state component of $I_p$ will affect the voltage trajectory most effectively when membrane conductance is low, which corresponds to the interburst interval. In trigeminal motoneurons, blocking this maintained outward current (i.e., the steady-state component of $I_p$) with cardiac glycosides shortens cycle time by effectively increasing the contribution of the inward currents during the interburst interval and thereby hastens burst onset. Sakai et al. (1996) similarly argue that the steady-state component of $I_p$ in rabbit sinoatrial node cells influences the pacemaker depolarization phase, the cardiac equivalent of the interburst interval. In addition to the steady-state component of $I_p$ that mostly influences the interburst interval, additional pump current activates specifically during the burst because of spike-mediated enhanced Na$^+$ influx. This dynamic component of $I_p$ constrains intraburst spike frequency and generates a transient postburst AHP (Fig. 6). We propose that the dynamic component of $I_p$ is responsible for burst termination in trigeminal motoneurons under apamin and low-Ca$^{2+}$/Ba$^{2+}$ conditions. Consequently, during the ouabain-induced transitional bursting phase, the intraburst $f$-$t$ relationship became erratic (and ultimately broke down), and the postburst AHP consistently declined in amplitude (Figs. 3 and 4). Once $I_p$ was fully blocked by the cardiac glycosides, every trigeminal motoneuron tested locked into the tonic spiking state that represents a nonterminating active phase.

A similar dynamic component of $I_p$ was characterized by Johnson et al. (1992) in rat midbrain dopaminergic neurons. Similar to our data, eliminating $I_p$ caused cessation of NMDA-induced bursting in midbrain dopaminergic neurons. These neurons, like trigeminal motoneurons, entered a nonterminating active phase of tonic spiking. Also similar to trigeminal motoneurons, the dynamic component of $I_p$ (the component of $I_p$ activated specifically during bursts) in dopaminergic neurons was sufficient to terminate bursts when $I_{\text{K,Ca}}$ was blocked pharmacologically (Johnson and Seutin 1997; Seutin et al. 1993). $I_p$ is also crucial for burst termination in several other neuron types (Angstadt and Friesen 1991; de Wael et al. 1993; Tsai and Chen 1995). In these cells, too, blocking $I_p$ induced tonic spiking.

Although burst termination does not critically depend on the slow inactivation of the persistent Na$^+$ and L-type Ca$^{2+}$ currents, these processes probably still occur and influence burst characteristics. We showed earlier that bath application of Bay K 8644 prolongs burst duration during serotonin-induced bursting (Hsiao et al. 1998), most likely by slowing the kinetics of the L-type Ca$^{2+}$ channels. This suggests that L-channel inactivation normally influences burst duration. Persistent Na$^+$ currents can also inactivate (Fleidervish et al. 1996), which could modify burst duration. Also during simulated bursts in this study (Fig. 6) the intraburst $f$-$t$ relationship still declined monotonically in the presence of apamin + ouabain. This suggests some temporal inactivation of the inward currents or recruitment of an unidentified outward current other than $I_{\text{K,Ca}}$ and $I_p$ during the simulated burst.

We propose that inactivation of the inward currents indirectly causes shorter burst duration during blockade of $I_p$. If the inward currents (L-type Ca$^{2+}$ or persistent Na$^+$ current) inactivate to some extent during bursts they normally de-inactivate during the relatively hyperpolarized interburst interval. Because removal of the tonic component of $I_p$ significantly shortens the interburst interval (described previously), the inward currents may not fully recover from inactivation between bursts. The magnitude of inward current available to generate subsequent bursts would then be compromised. These “compromised” bursts would then require less repolarizing current and therefore less time to terminate.

This proposal posits that less inward current underlies bursts after blockade of $I_p$. If this is true we would also expect an overall reduction in intraburst spike frequency after blockade of $I_p$, and specifically we would expect a downward shift in the intraburst $f$-$t$ curve. In fact, the intraburst $f$-$t$ curve in the presence of apamin + ouabain (Fig. 3C) was notably depressed compared with the intraburst $f$-$t$ curve during apamin alone (Fig. 3A). This suggests that the inward currents inactivated to some extent after application of cardiac glycosides, consequently depressing the intraburst $f$-$t$ curve and shortening burst duration.

In addition to blocking $I_p$, cardiac glycosides can induce “slip-mode conductance” in cardiac Na$^+$ channels, making them Ca$^{2+}$ impermeable (Santana et al. 1998). If this effect occurs in neuronal Na$^+$ channels it could potentially enhance inward currents during cardiac glycoside-challenged bursting and cause tonic spiking. However, we believe this effect cannot explain the current results because intraburst spike frequency...
clearly decreases during the ouabain and strophanthidin experiments (e.g., Fig. 3C). Decreased intraburst spike frequency is inconsistent with enhancement of inward current.

The tonic spiking state induced by cardiac glycosides is unlikely to represent general rundown in trigeminal motoneurons caused by large shifts in the Na⁺ and K⁺ concentration gradients for the following reasons. In the presence of apamin + strophanthidin bursts could initiate normally (Fig. 5), and the evoked action potential was close to full amplitude (Fig. 3B, bottom).

**Physiological significance**

Trigeminal motoneurons exhibit bistability and bursting in the presence of serotonin (Hsiao et al. 1998), an endogenous neuromessenger associated with oral-motor behaviors (Fornal et al. 1996; Hsiao et al. 1997; Veasey et al. 1995). Although rhythmic oral-motor behaviors are produced by brain stem central pattern generators (CPGs) that project to trigeminal motoneurons (Goldberg and Chandler 1990; Lund 1976; Nakamura and Kataoka 1995), the intrinsic properties of the motoneurons shape the final output pattern. In trigeminal (Hsiao et al. 1998) and spinal motoneurons (Hounsgaard et al. 1988a; Kieln 1991; Kieln and Eken 1997; Kieln et al. 1996) bistability and bursting are putatively important for motor behaviors; these properties can amplify motoneuronal output in the absence of maintained synaptic excitation. Because trigeminal motoneurons possess both I_{Ca,K} and I_p, and these two currents influence the temporal characteristics of serotonin-induced bursting, neuromodulation of I_{Ca,K} and I_p could regulate rhythmic motoneuronal spike output to match CPG activity. The regulation of I_p may simultaneously influence the phasic activity of the CPG because cardiac glycosides disrupt spinal networks that generate rhythmic motor patterns (Ballerini et al. 1997). In this way the intrinsic properties of trigeminal motoneurons would complement and assist the expression of ongoing spatiotemporal activity in the brain stem networks generating mastication or other rhythmic oral-motor behaviors, such as sucking, grooming, or licking.

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