Summation of Excitatory and Inhibitory Synaptic Inputs by Motoneurons With Highly Active Dendrites

Allison S. Hyngstrom, Michael D. Johnson, and C. J. Heckman

Department of Physiology and Department of Physical Medicine and Rehabilitation, Northwestern University, Chicago, Illinois

Submitted 13 November 2007; accepted in final form 26 January 2008

INTRODUCTION

Many types of neurons have voltage-sensitive conductances distributed throughout their dendritic trees (Gulledge et al. 2005; Hauser et al. 2000; Heckman et al. 2003; Magee and Johnston 2005; Migliore and Shepherd 2002). Spinal motoneurons have especially strong voltage-sensitive conductances in their dendrites (Hultborn et al. 2003; Lee and Heckman 2000; Prather et al. 2001). Excitatory and inhibitory synaptic inputs in motoneurons, as in other neurons, are largely located in dendritic regions (Fyffe 2001) and thus necessarily interact with dendritic voltage-sensitive channels before reaching the soma and generating spikes. Thus dendrites are active participants in synaptic integration, providing the potential not only for strong input amplification but also for highly nonlinear input interactions. Neuromodulatory input may also influence synaptic integration via the facilitation of voltage-sensitive conductances. These neuromodulatory inputs are likely to be tonically active in vivo (Aston-Jones et al. 2000; Jacobs et al. 2002; Steriade 2001). The effect of neuromodulators on voltage-sensitive conductances is not limited to the soma but extends to the dendrites (Frick and Johnston 2005; Heckman et al. 2003). Motor output is not generated solely by excitatory input but can also involve significant levels of inhibition (Berg et al. 2007; Perreault 2002). Thus normal synaptic integration in vivo depends on the interactions of neuromodulatory input with both excitation and inhibition.

In this study, we investigate the linearity of summation of excitatory and inhibitory synaptic inputs to the dendrites of adult spinal motoneurons subject to strong neuromodulation. We use a preparation of the feline spinal cord that provides two important advantages: selective activation of ionotropic synaptic inputs, via afferents from muscles, and tonic activity in a major neuromodulatory input from the brain stem, via axons releasing the monoamines serotonin and norepinephrine (Hounsagar et al. 1988). Our voltage-clamp technique prevents voltage-sensitive channels at the soma from influencing synaptic input, forcing the interactions between intrinsic and synaptic currents to occur in the relatively unclamped dendritic tree (Heckman and Lee 2001). Additionally, our technique also prevents synaptic input to the soma from affecting somatic voltage-sensitive channels. Voltage clamp also allows us to measure input summation as a function of voltage level at the soma to help identify sources of nonlinear interactions.

Previous studies using these methods have shown that monoaminergic input facilitates persistent inward currents (PICs) in motoneuron dendrites (Carlin et al. 2000; Hounsgaard and Kiehn 1993; Lee and Heckman 1996). Dendritic PICs amplify excitatory synaptic currents by as much as five- to tenfold (Hultborn et al. 2003; Lee and Heckman 2000; Lee et al. 2003; Prather et al. 2001). Once the cell is depolarized sufficiently to achieve full PIC amplification, a steep decline in the total excitatory current occurs in more depolarized membrane potential ranges (Lee and Heckman 2000) (see Fig. 1C). Nonetheless, two sources of excitation sum in a nearly linear fashion during repetitive firing (Prather et al. 2001). Inhibition effectively suppresses the dendritic PIC, but in a voltage range that is more depolarized than that for excitatory input. PIC suppression therefore overlaps with the range where excitatory current declines (Hyngstrom et al. 2007; Kuo et al. 2003). Our results show that nonlinear summation tended to compensate for the differences in PIC interactions with excitation and inhibition. Thus the net current due to the summation of excitation and inhibition was approximately linear, depending on input source. Overall, in cells with strong PICs, nonlinear summation of excitation and inhibition does occur, but this nonlinearity results in a more consistent relationship between membrane potential and the summed excitatory and inhibitory current.
and inhibition was more consistent (i.e., varied less) as a function of membrane voltage. Because the PIC has a major effect on repetitive firing in motoneurons, this compensation and resulting consistency across membrane voltage range is likely to play a significant role in determining motor output when the motoneuron is driven by combinations of excitation and inhibition.

**METHODS**

**Surgical preparation**

All surgical procedures were approved by the Northwestern University Institutional Animal Use and Care Committee. All experiments (total of 21) used the decerebrate cat preparation (for details see Lee and Heckman 1998b). The surgical preparation was completed under deep anesthesia (1.5–3% isoflurane in a 1:3 mixture of O2 and N2O). The right carotid artery was cannulated to monitor blood pressure and the right jugular vein was cannulated for delivery of drugs. The mixed nerves to the medial gastrocnemius and lateral gastrocnemius and soleus (MG and LGS) muscles were located and cuff electrodes were applied for antidromic identification of MG and LGS motoneurons (all experiments). Additionally, the common cutaneous sural nerve (5 experiments) and common peroneal (CP) nerve (12 experiments) were isolated and cuff electrodes applied for generating excitatory and inhibitory synaptic input to MG and LGS motoneurons. The MG and LGS (all experiments) and tibialis anterior/extensor digitorum longus tendons (TA/EDL, 11 experiments) were cut distally and secured to a tendon vibrator. A bilateral pneumothorax was performed to promote stable recordings by lessening movement of the chest wall. A laminectomy was completed from L1 to L7 vertebrae to expose the spinal cord and the cord was then bathed in mineral oil. Motor thresholds were determined for MG and LGS nerves. The animal was then paralyzed with Flaxedil (120 mg initial dose, then supplemented as needed) and ventilated. In 6 of the experiments, to diminish descending monoaminergic drive from brain stem centers, the cord was spinalized at T10 prior to decerebration. A laminectomy was done at T9,10 and two sutures were placed under the cord. The cord was then transected between the sutures. A precollicular decerebration was performed and all forebrain anterior to the lesion was aspirated and replaced with loosely packed saline-soaked cotton. The anesthesia was discontinued and recordings began 45 min after decerebration.

**Electrophysiology**

Intracellular recordings of lumbar motoneurons were done using sharp electrodes with resistances of 3–5 MΩ. Electrodes were filled with 2 M potassium citrate. Voltage-clamp recordings were performed using the Axoclamp 2A amplifier (Axon Instruments) in single-electrode discontinuous mode (9- to 10.5-kHz switching frequency). Due to the large current necessary to voltage clamp the motoneurons, an external feedback loop was added to increase the gain by a factor of 11 (Heckman and Lee 2001). This gain factor resulted in excellent clamp control, with deviations of <0.5 mV in voltage ramps. Voltage-clamp data were smoothed by digital filtering (~3 dB point of 0.3 kHz).

**Synaptic input**

Ia monosynaptic excitatory synaptic input to MG and LGS motoneurons was achieved by vibration of the Achilles tendon (180 Hz, 80 µm peak-to-peak). Muscle length was set to about 5 to 7 mm longer than slack length. This degree of stretch provides good driving of muscle spindle Ia afferents during vibration (Lee and Heckman 1996, 2000). In six experiments, polysynaptic excitatory synaptic input to MG motoneurons was generated by electrical stimulation of the caudal cutaneous sural nerve (100 Hz) (1.1–1.4 × threshold; this low intensity avoids the inhibitory component that can accompany sural input; sural input was not applied to LGS motoneurons; see LaBella et al. 1989). Reciprocal Ia inhibition to
MG and LGS motoneurons was produced by 100-Hz electrical stimulation of the CP nerve at 1.2–2.0 × motor threshold or by vibration of the common TA/EDL tendon (180 Hz, 80 μm peak-to-peak). The protocol used to assess these inputs (next section) assumes that they are constant for about 5 s. Although both Ia and sural excitation exhibit a slight decay in the first 1 s of steady stimulation, current thereafter is steady for many seconds (e.g., Lee and Heckman 2000; Prather et al. 2001). Many types of inhibition to motoneurons fade rapidly to repetitive stimulation (Heckman et al. 1994; Lafleur et al. 1993a, b); this is not true of the inputs used here, which thus meet the 5-s criterion (Kuo et al. 2003; Lee and Heckman 2000).

Protocol and data analysis

MG and LGS motoneurons were first identified by antidromic stimulation of peripheral nerves. In discontinuous voltage-clamp mode, the cell was held 5–10 mV hyperpolarized to its resting potential, after which a slow (6–8 mV/s) voltage ramp was applied in the absence of synaptic input. The same ramp was repeated in the presence of excitatory and inhibitory synaptic input alone and simultaneously. A second voltage ramp with no additional synaptic input was repeated to ensure cell health was maintained. Order of synaptic inputs was varied to prevent biased effects. Effective synaptic current ($I_s$) was calculated by subtracting the current traces from the synaptic trials from an average of control trials. Input conductance was determined by fitting a line to the hyperpolarized portion of the cell’s current–voltage ($I−V$) function. By subtracting the cell’s $I−V$ function from the line fitted to the hyperpolarized region we were able to determine PIC amplitude. Criteria for cell acceptance included: 1) antidromic spike height of $≥60$ mV; 2) holding current required to maintain the initial holding potential did not change $>20%$; 3) control input conductance did not change $>20%$; 4) the peak amplitude of the persistent inward current measured during control trials did not decline by $>20%$; and 5) adequate settling of the electrode during the recording. Consistency of the PIC was the main source of measurement error. To avoid systematic effects of this error, presentation order of excitation and inhibition was varied from cell to cell. Student’s $t$-test and paired $t$-test were used to make significance comparisons ($P < 0.05$ and ANCOVAs were used to compare differences in the slopes and intercepts between regression lines ($P < 0.05$). Excel and SPSS programs were used for data processing and data analysis.

RESULTS

Synaptic and intrinsic dendritic conductances shape the current–voltage function

Synaptic integration in spinal motoneurons was studied under two states of monoaminergic drive to the cord. In the medium monoaminergic state, the cord was intact and thus subject to tonic activity from descending monoaminergic inputs that is present in the decerebrate preparation (Hounsgaard et al. 1988) [The high monoaminergic state, achieved by exogenous administration of monoaminergic agonists (Lee and Heckman 1999), was not studied here.] In the low monoaminergic state, the cord was fully transected at the thoracic level to eliminate descending monoaminergic input (Hounsgaard et al. 1988). All studies were carried out in quasi–steady-state conditions, with both excitatory and inhibitory inputs applied as steady backgrounds. The basic protocol for all cells is illustrated in Fig. 1A, using Ia excitation and Ia reciprocal inhibition (see METHODS). The current–voltage function of the cell was assessed in control conditions (dotted line), with steady backgrounds of Ia excitation (dashed trace) and Ia inhibition (top thin trace). Results for simultaneous activation of both inputs are considered later.

The PIC is evident in the control $I−V$ function as a strong downward deflection, producing a negative slope region (see control trace, Fig. 1A) (Delgado-Lezama et al. 1997; Lee and Heckman 1996, 1998a; Schwinding and Crill 1980). The large changes in the $I−V$ function during the synaptic background conditions are due to interactions between the PIC and these synaptic inputs in dendritic regions, outside of the good voltage-clamp control applied at the soma (Heckman and Lee 2001). The asymmetry of the PIC interaction with excitation and inhibition noted in the Introduction is evident in Fig. 1A: excitation hyperpolarizes and broadens the activation of the PIC (bottom trace) (Lee and Heckman 2000), whereas inhibition suppresses the PIC amplitude without altering its activation voltage (Kuo et al. 2003). These differences in interaction of excitation and inhibition with the PIC are especially clear in leaky subtract $I−V$ functions (Fig. 1B).

All subsequent results are derived from these basic $I−V$ function protocols. In addition to Ia excitation and Ia inhibition (in 14 cells), we also used stimulation of the common cutaneous sural nerve alone ($n = 3$) and in combination with Ia excitation ($n = 3$). No significant differences were found for Ia versus sural excitation and these results were combined. We also studied Ia excitation combined with inhibition from electrical stimulation of a mixed nerve, the common peroneal (CP; $n = 11$). Both types of inhibitory inputs significantly increased input conductance ($P < 0.001$ in both cases, paired $t$-test), but the average percentage change in conductance for CP inhibition was significantly larger (Ia: $9 ± 10%$; CP: $35 ± 25%$; $P < 0.02$, $t$-test). Neither excitatory input significantly altered input conductance, which is consistent with previous work (Heckman and Binder 1988; Heckman et al. 1994). It should be emphasized that all sources of excitation and inhibition used in this study produce reasonably steady currents for the duration of the voltage ramps used here (see METHODS).

Amplification of excitatory and inhibitory inputs occurs in different voltage ranges

The effective synaptic ($I_N$) currents generated by Ia excitation and Ia reciprocal inhibition are shown in Fig. 1C. $I_N$ is the net current generated at the soma by a synaptic input (Heckman and Binder 1988, 1991) and can be calculated by subtracting the control $I−V$ function in Fig. 1A from each of the steady synaptic background $I−V$ functions (Lee and Heckman 2000). Because the ramps were applied slowly, this subtraction process closely approximates the $I_N$ that would be generated if the membrane potential was stepped to a steady holding potential, synaptic input applied, and the change in current measured (Lee and Heckman 1996, 2000). ($I_N$ was inverted in this study to make excitation positive and inhibition negative, simplifying presentation of summation results.) Applying both excitation and inhibition in the same cell produced results consistent with previous work in which these inputs were studied in different cell samples. The cell in Fig. 1C displays very marked amplification of excitation (+Ia $I_N$; Fig. 1C, dashed trace), with a peak around $−45$ mV, followed by strong reduction above $−45$ mV (cf. Lee and Heckman 2000). This strong decline in excitatory
Summation of excitation and inhibition during dendritic PIC amplification

In a cell with perfect voltage and space clamp, the current generated by ionotropic synaptic inputs varies linearly with voltage and also undergoes linear summation. This linearity occurs because voltage-clamp controls the driving force for the entire membrane of the cell. In the present study, only the soma is well clamped. Therefore the amplification and saturation of synaptic input due to the dendritic PIC (Fig. 1C) may also induce nonlinear input summation in dendritic regions. In other words, the good voltage clamp at the soma (see METHODS) provides the advantage of focusing our results on the nonlinear interactions between synaptic input and the PIC in dendritic regions.

Linearity of summation was assessed by comparing the actual summed $I_{S_N}$ generated by simultaneous activation of both excitation and inhibition, to the summed $I_{S_N}$ calculated from linear addition of the $I_{S_N}$ currents generated by independent activation of excitation and inhibition (e.g., Powers and Binder 2000). Average $I_{S_N}$ was determined for three 5 mV voltage windows (sub-PIC, peak excitatory $I_{S_N}$, and peak inhibitory $I_{S_N}$; see Fig. 2C). Thus nonlinear summation is defined by deviations of actual from linear. Examples are shown in Fig. 2. The linear summed $I_{S_N}$ currents (dotted traces) exhibited large variations in amplitude as holding potential was depolarized, with net excitation tending to dominate near the voltage for peak excitatory amplification and net inhibition at the more depolarized voltage for peak inhibition. The actual summed $I_{S_N}$ currents (thick traces) exhibit deviations from the linear summed $I_{S_N}$ currents (dotted traces), indicating the presence of nonlinear summation. It is evident that this nonlinear summation results in deviations both above and below the linear prediction. We have adopted the following terminology to describe these deviations. In the voltage range where peak amplification of excitatory input occurred, actual summed $I_{S_N}$ was less positive than the linear summed $I_{S_N}$, giving sublinear summation. In contrast, at the more depolarized voltage range where peak amplification of inhibition occurred, the actual summed $I_{S_N}$ was more positive than the linear summed $I_{S_N}$, giving supralinear summation. The examples in Fig. 2 show the full range of nonlinear behavior, from large (Fig. 2, A and C) to moderate (Fig. 2B).

The trend for a transition from sub- to supralinear summation was consistent in the data set as a whole. Summation was quantified in three voltage windows, each 5 mV wide (see the horizontal lines in the examples in Fig. 2). Figure 3 plots actual versus predicted results for each neuron in each window. Figure 3A shows that, in the sub-PIC voltage window (centered 5–10 mV hyperpolarized to the resting potential), sublinear summation occurred in many cells, but overall this trend was not significant (actual $I_{S_N}$ differed from predicted $I_{S_N}$ by $-0.6 \pm 1.7$ nA; $P = 0.064$, paired $t$-test). For the voltage window centered on the peak of excitatory amplification (Fig. 3B), actual $I_{S_N}$ tended to be sublinear by an average of $-1.5 \pm 2.7$ nA and this difference was significant ($P < 0.003$, paired $t$-test). In contrast, within the voltage window centered on the peak of inhibitory amplification (Fig. 3C), summation was significantly supralinear (actual more positive than predicted) (difference of $+1.9 \pm 3.4$ nA; $P < 0.003$, paired $t$-test). The relative nonlinearity in each voltage window was calculated by dividing these average nonlinearities by the absolute value of the sum of the applied inputs. This procedure gave equal weight to both excitatory and inhibitory inputs. In the peak excitatory and inhibitory windows, nonlinear summation was found to be of similar magnitude, averaging 15% sublinear in the peak excitatory voltage window and 18% supralinear in the peak inhibitory window. Each of these average percentages is significantly different from zero ($t$-test, $P = 0.0004$ and $P = 0.0007$, respectively). In the sub-PIC window, the average of 6% was not significantly different from zero.

Figure 3 also shows that, in all voltage windows, the input combinations spanned a wide range, from inhibition being dominant (lower left quadrants) to excitation being dominant (upper right quadrants) (each data point is from a different cell). Thus our studies were not biased by a narrow range of relative amplitudes of excitation and inhibition. The amplitude of nonlinearity for each cell in Fig. 3 is indicated by the vertical distance from the line where $y = x$ (thick line). There was no tendency for nonlinear summation to increase or decrease with relative amount of inhibition versus excitation and thus linear regression satisfactorily fit data for all voltage windows. The deviations from linearity were not significantly correlated with basic motoneuron properties (e.g., input conductance) or with PIC parameters (e.g., amplitude, voltage threshold). In all three windows, the slopes of the regression relations were not significantly different from 1.0 (i.e., the 95% confidence intervals for these slopes included the $y = x$ line). We also divided the data sets according to source of input. We found no significant differences in slope or intercept for the regression relations for the data in Fig. 3 in cells where CP stimulation was the source of inhibition compared with cells with Ia inhibition, nor when sural excitation was used instead of Ia excitation (ANCOVA, $P > 0.05$ in all cases).

A striking feature of the data was the transition from sub- to supralinear summation as holding potential was depolarized from the voltage window for peak excitation to peak inhibition. To quantify the net impact of this transition, we calculated the difference between summed $I_{S_N}$ at the voltages for peak amplification of excitation and for peak amplification of inhibition ($\Delta$Sum; see arrows in Fig. 2B). Overall, the transition from sub- to supralinear summation reduced $\Delta$Sum from 7.7 ± 4.9 nA for the linear summation case to 4.1 ± 2.9 nA in the actual case ($P < 0.0001$; for examples, see Fig. 2). Thus nonlinear summation reduced the difference in $I_{S_N}$ between the voltage windows for peak excitatory amplification and peak inhibitory...
amplification. Further analysis revealed that when the linear ΔSum was large (as for the cell in Fig. 2A), the nonlinear summation increased, resulting in a relatively larger reduction in actual ΔSum (again, see Fig. 2A). To illustrate this result quantitatively, Fig. 4 plots the difference between linear and actual ΔSum as a function of the baseline of linear ΔSum. A strong relationship is apparent ($r = 0.81$; $r^2 = 0.65$; $P < 0.001$). For large values of predicted ΔSum, the reduction in actual ΔSum due to nonlinear summation was substantial, as much as 10 to 15 nA (Fig. 4). Changes of these magnitudes are certainly large enough to be functionally relevant during repetitive firing (see DISCUSSION).
Summation behavior when dendritic PIC amplification is suppressed due to lack of monoaminergic input

To allow summation to be assessed without PICs, we studied cells in a preparation with minimal monoaminergic drive (acutely spinalized decerebrate; see METHODS). PIC amplitudes in these cells were found to be small [mean PIC amplitude = −3.6 ± 3.1 nA, n = 18; for the cells in the medium monoaminergic state in the previous section, mean PIC amplitude was −12.2 ± 6.4 nA; this difference is consistent with our previous studies (Kuo et al. 2003; Lee and Heckman 2000)]. The I–V functions for cells in the minimal monoaminergic state were much more linear than in the medium state, as illustrated by the example in Fig. 5A, and neither excitatory nor inhibitory \( I_N \) currents, summation behavior was calculated in only two 5-mV-wide windows. The hyperpolarized voltage window was set 5–10 mV hyperpolarized to rest (which was similar to the sub-PIC window in the medium state). The depolarized voltage window was centered where the small PIC reached its peak value following leak subtraction. In cells with no clear PIC (as in Fig. 5), the depolarized window was placed 10–15 mV above the hyperpolarized window because this was the typical position of the PIC relative to the rest level.

Excitatory \( I_a \) \( I_N \) was small in both voltage windows (hyperpolarized window: 2.8 ± 2.2 nA; depolarized window: 4.0 ± 3.0 nA) and was not significantly different from the mean excitatory \( I_N \) measured in the sub-PIC voltage window in the medium monoaminergic state (cord intact) (\( P > 0.24 \)). Both \( I_a \) and CP inhibition remained similar in intensity compared with the medium monoaminergic state because percentage changes in input conductance in the two states were not significantly different from each other (\( t \)-test, \( P > 0.05 \)).

With respect to the sub-PIC voltage window in the medium monoaminergic state (Fig. 3A), summation in the hyperpolarized window in the low monoaminergic state tended to be approximately linear (square symbols, Fig. 6) (difference between observed and predicted: −0.42 ± 2.73 nA, \( P > 0.5 \)). In the depolarized window (triangle symbols), summation tended to be slightly sublinear (difference of −1.68 ± 2.86 nA, \( P < 0.03 \)), much as for the peak excitatory voltage window in the medium monoaminergic state (Fig. 3B). In both windows, the cells where CP stimulation was the source of inhibition tended to behave differently from the cells where \( I_a \) reciprocal inhibition was combined with excitation. For CP inhibition, the

![Fig. 3](image-url)

**FIG. 3.** Plots of actual vs. predicted summed \( I_N \) currents for each cell in each voltage window. The actual summation of excitatory and inhibitory \( I_N \) currents was plotted against the predicted \( I_N \) for 3 voltage windows in the medium monoaminergic state: A: sub-PIC window (\( y = 0.76x - 0.4 \)); B: peak excitatory window (\( y = 0.90x - 1.3 \)); C: peak inhibitory window (\( y = 0.92x + 1.74 \)). In each plot the thick line is a hypothetical case of linear summation where \( y = x \). Vertical distance from this line represents nonlinear summation. The solid lines indicate the regression relations and the dashed lines the 95% confidence limits for the slopes.

![Fig. 4](image-url)

**FIG. 4.** When linear addition predicts that summed \( I_N \) exhibits a large variation as membrane potential is depolarized, compensation for this variation by nonlinear summation increases. In other words, when the \( \Delta \)Sum parameter (defined in Fig. 2B) for linear summation is large, nonlinear summation markedly reduces actual \( \Delta \)Sum (as in the cell in Fig. 2A). This compensation is illustrated quantitatively by the plot shown here, where the y-axis is the reduction in \( \Delta \)Sum (linear \( \Delta \)Sum − actual \( \Delta \)Sum) and the x-axis is the baseline of linear \( \Delta \)Sum (y-axis) (\( r = 0.81, r^2 = 0.65; P < 0.001 \)). This relationship is functionally relevant in that the reduction in \( \Delta \)Sum reaches currents of 10 to 15 nA (top right), which would modulate firing by as much as 30 Hz (cf. Lee et al. 2003).
95% confidence intervals of the slopes of the regression relations included the line \( y = x \) (both windows). In contrast, regression slopes for the data with Ia inhibition were shallow and significantly <1.0 (slope = 0.39 in the subthreshold window; slope = 0.34 in the peak PIC window; in both cases the 95% confidence intervals did not include slope = 1.0). This Ia versus CP difference may be a consequence of the relative balance of excitation and inhibition. As noted in the previous section, CP inhibition tended to be stronger than Ia inhibition and thus the CP data points tend to lie in the lower left quadrant of Fig. 6, where inhibition is greater than excitation.

D I S C U S S I O N

The dendritic PIC has a tremendous influence on synaptic integration in motoneurons (Binder 2002; Heckman et al. 2003; Hultborn et al. 2004). Because the PIC amplification factors are two- to threefold, the dendritic PIC generates more net current at the soma (i.e., more \( I_C \)) than does the synaptic input on its own. Excitation on its own and inhibition on its own both tend to interact in a proportional manner with the PIC (Hultborn et al. 2003; Kuo et al. 2003; Prather et al. 2001). Our results revealed, however, that simultaneous activation of excitation and inhibition induced significant nonlinear summation. Furthermore, this nonlinear interaction had a very distinct form, transitioning from sublinear to supralinear as voltage-clamp holding potential was progressively depolarized. This transition compensated for an important difference in the interaction of the dendritic PIC with excitation compared with inhibition. Excitation tended to undergo maximum PIC amplification in a voltage range at which spikes would be initiated in unclamped conditions, whereas inhibition tended to be amplified at significantly more depolarized levels. The interaction between excitation and inhibition is thus inherently unbalanced, with excitation dominating in the region near spike threshold and inhibition just above (see predicted summed currents in Fig. 2, A–C). Nonlinear summation tended to restore a more consistent relation between these two inputs as membrane potential was depolarized, with this restoring effect increasing in proportion to the potential imbalance (see Fig. 4).

Possible mechanisms for nonlinear summation in the medium monoaminergic state

An important concern with any study of summation of synaptic inputs is that the input interactions may occur partially by presynaptic mechanisms. For motoneurons, interactions could occur among premotor interneurons, which are activated by the sensory afferents, or by presynaptic inhibition of afferent terminals, which is a fundamental feature of cord circuitry (Rudomin 2002). The inputs we used were selected to minimize these problems. The Ia excitation is primarily monosyn-
aptic. A small multisynaptic component may exist, but experiments in our lab have yet to detect any effects not due to ionotropic, monosynaptic excitation (Kuo et al. 2003; Lee and Heckman 2000). Note also that the lack of amplification of Ia \( I_{H} \) in the low monoaminergic state indicates that this input does not significantly influence dendritic PICs by either metabotropic or N-methyl-D-aspartate glutamate receptors (see also Heckman et al. 2000). Ia afferents are subject to presynaptic inhibition and the cutaneous afferents activated via the electrical stimulation of the CP nerve could be a significant source of this form of inhibition (Baldissera et al. 1981). Nonetheless, presynaptic inhibition of afferents should reduce Ia and sural excitation and thus appear as a strong sublinear summation. In contrast, CP stimulation generally produced linear summation in the sub-PIC range in the low and medium monoaminergic states. Moreover, in the medium monoaminergic state, the transition from sub- to supralinear summation in the depolarized voltage windows (peak excitation and peak inhibition) indicates that presynaptic inhibition, if present, was not a dominant effect. These findings do not eliminate a role for presynaptic inhibition in our results, but do suggest that it was not large compared with other effects, such as those due to voltage-sensitive conductances. Consequently, it is likely that the dominant interactions determining linearity of summation occurred postsynaptically. Finally, it should be noted that Moritz et al. (2007) recently showed that PICs in the somatic region of motoneurons are strong and could thus amplify input (cf. Stuart and Sakmann 1995). The secure voltage clamp of the soma in the present studies, however, restricts amplification, saturation, and nonlinear interactions to regions of poor voltage control (i.e., the dendrites).

Inhibition tends to suppress the dendritic PIC (Fig. 1; Kuo et al. 2003) and thus should reduce the amplification of excitatory input. This suppression could account for sublinear summation in the voltage range for the peak excitatory \( I_{H} \) in the medium monoaminergic state. Further depolarization brings on saturation of the Ia excitation when applied on its own (Fig. 1B), presumably due to dendritic depolarization or perhaps activation of voltage-sensitive outward currents. Simultaneous dendritic inhibition could provide hyperpolarization to reduce this saturation (Fig. 2) and thus result in supralinear summation.

Possible mechanisms underlying summation with minimal PIC activation

In the sub-PIC voltage window where hyperpolarization minimized the activation of the dendritic PIC by synaptic input, other voltage-sensitive conductances may still influence synaptic integration. In particular, the hyperpolarization-activated cationic current \( I_{H} \), which is responsible for the “sag” to hyperpolarizing steps in motoneurons (Powers and Binder 2001), may play a significant role. Activation of \( I_{H} \) might offset the shunting effect of inhibition and account for the nearly linear summation seen in Fig. 3A. In the present study, such an interaction would have taken place in the poorly clamped dendritic regions. This interpretation would therefore imply that motoneurons have substantial \( I_{H} \) in dendritic regions, a possibility not yet investigated for motoneurons via immunohistochemical methods but supported by the experimental and modeling studies of Manuel et al. (2007).

In the low monoaminergic state, summation behavior in the hyperpolarized window was similar to that in the sub-PIC window in the medium state (i.e., approximately linear). The reason for the difference between behaviors with CP versus Ia inhibition are not clear, but dendritic \( I_{H} \) may again be involved. Note that in Fig. 6, the low slope for the regression relation for the Ia inhibition means that this data set tends to merge into the CP data as inhibition becomes large compared with excitation (moving toward the lower left on this plot). This trend is consistent with larger relative amounts of inhibition generating input mixtures more likely to active \( I_{H} \) in dendritic regions. This explanation may not be sufficient, however, in that the behavior of Ia inhibition was the same in both the hyperpolarized and depolarized voltage windows. The PIC was small enough in these low-monoaminergic-state data not to have much influence at the depolarized level, but the current injected for the somatic voltage clamp might have pushed the dendritic regions above the range of \( I_{H} \) activation. Further study of the effect of \( I_{H} \) on combinations of excitation and inhibition is warranted, especially in light of recent work with dynamic clamp showing that \( I_{H} \) can produce a resonant peak in synaptic current (Manuel et al. 2007).

Functional relevance of nonlinear summation of excitation and inhibition in normal motor behavior

The degree to which inhibition is actually mixed with excitation during normal motor behavior is not presently known. Berg et al. (2007) demonstrated clear comodulation of excitation and inhibition during the scratch reflex in the turtle. Although inhibition of this magnitude may not be present in other motor behaviors (see Perreault 2002), chronic recordings of spinal interneurons in the primate have shown that most of these presumably premotor cells are tonically active even in the resting state (Prut and Perlmutter 2003). These studies included interneurons with inhibitory actions on motoneurons. Thus significant inhibition may often be mixed with excitation during motor output.

Motor output is of course generated by repetitive firing in motoneurons. The voltage-clamp method used in this study had the advantage of restricting nonlinear interactions between the inputs to unclamped dendritic regions (Heckman and Lee 2001). These results are thus likely to provide key data for constraining computer simulations of dendritic processing in motoneurons, which, given the near impossibility of direct dendritic intracellular recordings in motoneurons, continue to be essential for understanding the dendritic PIC (Elbasiouny et al. 2005, 2006; Grande et al. 2007a,b). Yet our results may also be relevant to synaptic integration during repetitive firing (e.g., Prather et al. 2001, 2002). Our results in the minimal monoaminergic state showing sublinear to linear summation of excitation and inhibition during voltage clamp are consistent with those of Powers and Binder (2000) during repetitive firing in a deeply anesthetized preparation, likely to have similarly low levels of monoaminergic drive. These similarities between firing and voltage-clamp behaviors are reasonable in that the afterhyperpolarization (AHP) that follows each spike limits the average membrane depolarization between spikes (Powers and Binder 2001). The average voltage level during firing does depolarize substantially as firing rate increases (Schwindt and Calvin 1972; Schwindt and Crill 1982), reaching the levels...
where excitatory saturation and inhibitory amplification occur (Lee and Heckman 1998a,b, 2000). Thus it is probable that the transition from sub- to supralinear summation seen here in voltage clamp could also produce a more consistent balance between excitation and inhibition as baseline firing rate increases from threshold levels. Consider, for example, a case where perfect linear summation prevails and where a cell is firing at a low rate, near threshold, due to a mixture of excitation and inhibition (excitation dominating, of course, or firing would not occur). As the excitatory input is increased, average membrane potential becomes depolarized, pushing the cell into the region where inhibitory amplification is greater than excitatory amplification (the peak inhibitory windows in Fig. 2). This depolarization would increase the efficacy of inhibition already present and reduce the efficacy of the added excitation, very much limiting the resulting change in firing. This limiting of the effect of excitation would be exacerbated by the decline in amplification of excitatory $I_E$. The nonlinear summation process, with its transition from sub- to supralinear summation, would reduce these variations in excitatory versus inhibitory synaptic efficacy and allow firing rate to increase more in proportion to the increase in net input. In general, the extra noise induced by mixing inhibition with excitation may not be a significant functional problem because the large AHP in motoneurons acts to stabilize firing rate to noisy input (Manuel et al. 2005, 2006). Overall, the processing of excitation and inhibition in motoneurons with strong dendritic PIC's does involve nonlinear interactions although, to a significant degree, these nonlinearities tend to make the PIC processing of the input mixture more consistent as a function of membrane voltage.

ACKNOWLEDGMENTS

We thank J. Miller for assistance in data collection, J. Schuster for assistance in preparation of the manuscript, and J. Kuo for assistance with data collection and technical consultation.

GRANTS

This work was supported by National Institute of Neurological Disorders and Stroke Grant NS-034382 to C. J. Heckman; and U.S. National Research Service Award, Predoctoral Fellowship 5F31NS048757-03, and the Foundation for Physical Therapy, Promotion of Doctoral Studies, Viva J. Erickson Scholarship to A. S. Hyngstrom.

REFERENCES


Hultborn H, Brownstone RB, Toth TI, Gossard JP. Key mechanisms for setting the input-output gain across the motoneuron pool. Prog Brain Res 143: 77–95, 2004.


Rudomin P. Central control of information transmission through the intraspinal arborizations of sensory fibers examined 100 years after Ramon y Cajal. Prog Brain Res 136: 409–421, 2002.


