Intense Synaptic Activity Enhances Temporal Resolution in Spinal Motoneurons

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Abstract

In neurons, spike timing is determined by integration of synaptic potentials in delicate concert with intrinsic properties. Although the integration time is functionally crucial, it remains elusive during network activity. While mechanisms of rapid processing are well documented in sensory systems, agility in motor systems has received little attention. Here we analyze how intense synaptic activity affects integration time in spinal motoneurons during functional motor activity and report a 10-fold decrease. As a result, action potentials can only be predicted from the membrane potential within 10 ms of their occurrence and detected for less than 10 ms after their occurrence. Being shorter than the average inter-spike interval, the AHP has little effect on integration time and spike timing, which instead is entirely determined by fluctuations in membrane potential caused by the barrage of inhibitory and excitatory synaptic activity. By shortening the effective integration time, this intense synaptic input may serve to facilitate the generation of rapid changes in movements.

Introduction

Spike timing in nerve cells is determined by temporal integration of synaptic potentials and intrinsic response properties. However, little is known about the timescale of this integration during functional network activity and how it is affected by synaptic events. In the absence of synaptic input the spike afterhyperpolarization (AHP) determines spike timing during repetitive firing. In motoneurons (MNs), the frequency range of this firing is well suited for force regulation in the muscle fibers they innervate [1]. This suggests that main role of AHP is temporal filtering that converts the continuous asynchronous synaptic bombardment to a regular output discharge of action potentials. Furthermore, firing maintained by AHP and other slow intrinsic properties is also appealing because it is a metabolically inexpensive way of shaping the spike patterns to suit particular functions, e.g. spinal motor rhythms [2–5].

On the other hand, the AHP and other slow intrinsic properties would impede rapidly changing motor responses and it is not known how resilient they are to a noisy background of synaptic activity. Recent evidence suggests that intrinsic response properties may be shunted by synaptic conductance in cortical and subcortical networks [6–10]. In the spinal cord of the adult turtle, scratch motor network activity is associated with a dramatic rise in conductance and in fluctuations of the membrane potential (V_m) in both MNs and interneurons during spiking [11,12]. This is due to a concurrent intense inhibitory and excitatory synaptic activity [13]. Under these conditions of high synaptic conductance, the temporal resolution is predicted to be enhanced [14–16] and the role of slow intrinsic properties becomes less obvious. Surprisingly few experimental studies have explored this interplay between high synaptic conductance, AHP and temporal integration in active networks.

For this reason, we have conducted experiments on spinal motoneurons embedded in a functionally active network during fictive motor behavior. In earlier studies the conductance increase in motoneurons during fictive locomotor and scratch network activity was first measured in vivo in the cat [17,18] and in the turtle [12]. The isolated spinal cord-carapace preparation from the turtle [19] offers uniquely stable recording conditions in which intrinsic and synaptic conductance changes during network activity can be measured against a background of very low leak conductance [11,13,20]. This allows us for the first time to quantify the relative importance of active and passive intrinsic properties and the dynamics of synaptic input for spike timing during functional network activity.

We measured the effective integration time in MNs during network activity using a novel statistical approach that quantified the V_m-fluctuations before and after the action potential. Three temporal features were characterized: the membrane time-constant, effective synaptic integration time (eSIT) and the effective recovery time (eRT). We define the eSIT as the time it takes to sum up enough synaptic input to cause a spike. The eRT is defined as the time it takes for the V_m-distribution following a spike to return to the pre-spike condition, i.e. how long it takes the cell to “forget” that a spike has occurred. We report eRT as short as 4 ms during network activity, which is more than a 10-fold decrease compared with quiescent network. Our results show that even prominent intrinsic response properties like the AHP are severely attenuated concurrent with increase in synaptic conductance. For this reason, the contribution of synaptic activity and active membrane properties to network dynamics can only be captured by a
conductance-based model [8,16,21,22] (Figure 1). We conclude that a shortening of the recovery time and integration time of motoneurons cause an increase in the temporal resolution of the motor system during activity, which we suggest as a mechanism to facilitate rapidly changing movements.

Results

The question we ask is how the surge in conductance from intense synaptic bombardment during network activity modifies intrinsic properties exemplified by the spike AHP and shrinks response time and integration time of the neuron. The sensitivity of spike generation and spike pattern to conductance is illustrated in simulations in a simple model (Figure 1). In the absence of synaptic input the regular firing during maintained depolarization in the model neuron was determined by the AHP (red trace in Figure 1A). Synaptic input simulated as current noise, i.e. linear summation of synaptic potentials, resulted in greater variability in spike timing, but did not change the averaged AHP (black trace Figure 1A, B). Thus, current-based synaptic input fluctuations changed neither total conductance nor neuronal integration time. However, when synaptic activity was modeled more realistically as a noisy conductance increase, spike timing was strongly influenced by the stochastic fluctuations in $V_{m}$ (blue trace, Figure 1A, B), the AHP was severely attenuated (arrow in Figure 1B) and the integration time was shortened [22,23]. Thus, the simple simulation showed that a fluctuating current is not likely to affect the integration time or reduce the importance of the AHP whereas intense and fluctuating synaptic conductance is.

For the purpose of making these statements applicable to quantitative evaluation of experimental data, we developed statistics to capture the temporal features of $V_{m}$ before and after the spike. We define two measures linked to the superposition of spikes (Figure 1C and D). First, the effective synaptic integration time (eSIT), defined as the time period prior to spikes during which fluctuations in membrane potential showed a significant depolarizing trend compared with a baseline template (Figure 1D). Likewise, we define the effective recovery time (eRT), as the time it takes to regain pre-spike $V_{m}$-levels. Notice that both eRT and eSIT are absent in current noise model whereas they are present in the conductance noise model (cf. Figure 1C and D).

To evaluate whether the neuronal integration time and AHP are substantially affected during intense network activity we measured eSIT and eRT in MN during scratching. The data set consisted of more than 10,000 spikes in 185 scratch epochs in 17 MNs. The analysis is organized as follows. First, we estimated the conductance increase in a sub-sample of MN to confirm that strong synaptic components were present (Figure 2). Next, we applied our statistics to measure the eSIT and eRT and estimated the effective membrane time-constant from $V_{m}$-fluctuations at different levels of synaptic activity. Since intensity of synaptic activity may vary among cells, we compared the eRT with indicators of input, i.e. the effective membrane time constant as well as the smallest inter-spike interval (ISI) for each MN. Finally, since the functional expression of AHP accumulation in motoneurons is spike frequency adaptation (SFA) [4,24], we also tested for adaptation during each cycle of the scratch epochs.

Synaptic conductance

A scratch epoch was induced by rhythmic cutaneous stimulation of the skin in the hind-limb pocket and the concurrent synaptic conductance (Figure 2A, B) was estimated from the voltage deflections to injected current pulses (Figure 2C–F) for a subset of MN. Conductance due to action potentials is a potential source of spikes (Figure 1C and D). A template distribution of $V_{m}$ traces is chosen at an arbitrary time prior to the spike (see arrow at $t_{s}$, arrow) for which the distribution at the rest of the time points is compared with. The outcome of comparison is shown as the KS-test trace below, 1 represents acceptance and 0 represents rejection of the hypothesis that they are different. Below is shown the P-values for the KS test. The distribution of $V_{m}$ in (C) is different everywhere, whereas the distribution in (D) is only different up to and immediately after the spike. The time it takes to regain pre-spike distribution following the spike is referred to as effective synaptic integration time (eSIT, arrow). The estimated conductance increased 2–5 times during
We attribute this increase to balanced inhibitory and excitatory synaptic input [13] and conclude that temporal processing in MNs could be affected by synaptic conductance during network activity.

Effective synaptic integration time

We first considered the $V_m$-statistics prior to spikes. The time of the earliest statistical sign of depolarization prior to action potentials we dub “the effective synaptic integration time” (eSIT). This depolarization may be caused by a rise in excitatory conductance or a fall in inhibitory conductance. For each cell, the eSIT was estimated by comparing a template distribution of $V_m$ with distributions of $V_m$ as a function of time prior to action potentials, as in figure 1. The comparison was attained using a KS-two-sample test at each point in time (method, Figure 3). The template distribution was chosen at a time, $t_{\text{template}}$, well before the spikes (arrows Figures 3A and B). The eSIT did not depend systematically on the choice of $t_{\text{template}}$ as long as it was at least 10 ms prior to the spike (Figure 3C) even though the number of samples in the distribution decreased (Figure 3D).

Memory of a spike–eRT

The same statistical test was used to evaluate the impact of a spike by comparing the $V_m$-distribution after action potentials with template distributions well before the spike (arrow Figure 3B). In the graph of the KS-test, the gray area represents rejection of the null hypothesis that the $V_m$-distribution before and after the spike was statistically similar to the template distribution. When the $V_m$-distribution after the spike was indistinguishable from the template distribution, we considered the impact of a spike to have ceased (graph of $P$-values, Figures 3A and B). Both eRT and eSIT had no clear dependence on $t_{\text{template}}$ earlier than ~20 ms (Figure 3C). Since the number of spikes in the distribution decreased with $t_{\text{template}}$ (Figure 3D) the eRT estimate should not be performed with a high $t_{\text{template}}$. We chose to calculate the mean eRT over the range 20 ms $< t_{\text{template}} <$ 40 ms.

Impact of $\tau_{\text{eff}}$ and AHP on recovery time

Right after the occurrence of an action potential $V_m$ is hyperpolarized. The time it takes for $V_m$ to re-polarize back to the level prior to the spike depends on the AHP and on the passive effective time-constant of the membrane, which we refer to as $\tau_{\text{eff}}$. 

Table 1. Total input conductance for 5 MN during both quiescence and active network.

| Motoneuron number | $G_{\text{quiescence}}$ [nS] | $G_{\text{active}}$ [nS] | Fraction [%] |
|-------------------|--------------------------|--------------------------|--------------|
| 2                 | 73                       | 320                      | 438          |
| 3                 | 45                       | 181                      | 405          |
| 10                | 71                       | 224                      | 314          |
| 13                | 20                       | 40                       | 200          |
| 14                | 55                       | 258                      | 470          |

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Figure 2. Increase in synaptic conductance during a single scratch epoch. (A) Cutaneous stimulation via sinusoidal movements of a glass rod on the hind-limb pocket skin. (B) Electro-neurogram from hip flexor nerve. (C) $V_m$ from intracellular recording. Tick mark indicates $V_m$ of -100 mV. (D) Current pulses of 0.7 nA from constant current level of 1.0 nA. (E) High-pass filtered $V_m$ from (C) (cut off is 2 Hz). Transient artifacts removed. (F) left, average voltage deflections from (E) during quiescence (n = 14), right during network activity (n = 14). The average increase in conductance is 340%. Notice the occurrence of two spikes. (G) The membrane conductance as a function of time. The peak conductance during network activity is >800% of the conductance during quiescence.

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In our attempt to sort out which part of the re-polarization is due to the passive decay of $V_m$ and which is due to the cessation of AHP conductance, it is important to estimate $\tau_{ef}$ during network activity. The increase in total conductance will also diminish the relative importance of the AHP conductance and the AHP will appear shorter. To quantify the vestige of the AHP under different levels of input conductance we define the effective After-Hyperpolarisation Period ($eAHP$). This period represents the time it takes until the AHP ceases to affect the $V_m$ trajectory. If there is no overlap between the passive membrane decay and the $eAHP$, the $eRT$ is just the sum of $\tau_{ef}$ and $eAHP$, while if there is overlap $eRT$ will be less than the sum:

$$eRT \leq eAHP + \tau_{ef}.$$  

It is important to emphasize that $eRT$ also represents the upper bound on both $eAHP$ and $\tau_{ef}$, i.e. $eRT \geq eAHP$ and $eRT \geq \tau_{ef}$. Hence, it was important to determine $\tau_{ef}$ for each cell in order to determine the contribution of AHP to the integration process.

**Estimating $\tau_{ef}$ during activity**

For the spinal motor activity we differentiated three situations: The quiescent state with little or no synaptic input; the on-cycle with motor nerve activity and spike activity in motoneurons; and the off-cycle, which is at the low point in between the on-cycles (Figure 4A–C). As expected the effective membrane time constant decreased significantly during scratching. For representative data, $\tau_{ef}$ was 27 ms in the quiescent state, while it was only 2.8 ms in the On-cycle and 5.2 ms in the off-cycle (Figure 4C–E). Membrane time constants during quiescence are ordinarily estimated from $V_m$-decay times after injected current pulses (Figure S2). During the dynamic and intense synaptic input in the on-cycle and off-cycle, this method was both difficult and imprecise. Instead, if $V_m$ is assumed to follow a stochastic process known as Ornstein-Uhlenbeck-process (OU-process), then maximum likelihood estimation would be the proper way to obtain $\tau_{ef}$ [26,27]. The estimated values of $\tau_{ef}$ with this technique are listed in Data and Methods 1 in Data and Methods S1 and see Figures S1–S3. It turned out that $V_m$ did not obey an OU-process (Figure S4) and these estimates of $\tau_{ef}$ were systematically much higher than $\tau_{ef}$ and $eRT$. Therefore, $\tau_{ef}$ was instead estimated empirically by fitting an exponential decay to the initial part of the auto-correlation sequence (Figure 4F). It was necessary to fit to the initial part (from 0 to 3 ms, vertical gray line) to assure that the auto-correlation lag was small compared to the total length of the sample (200 ms), since the premise of exponential decay is infinite-length of data trace (see Data and Method S1).

**Temporal features for the population**

The effective synaptic integration times, the effective recovery times and the effective membrane time constants across the population of neurons are listed in Figure 5A. The average $eSIT$ was $\mu = 7.7 \pm 0.8$ ms and the average $eRT$ was $\mu = 7.5 \pm 0.7$ ms (mean$\pm$SE, n = 17). The median of $eSIT$ and $eRT$ were both 7.7 ms. The $eRT$ was assumed to be approximately equal to the sum of $\tau_{ef}$ and the effective AHP (see above). However, the population average of $\tau_{ef}$ was $\mu = 9.3 \pm 1.4$ ms (mean$\pm$SE, n = 17) while the median of $\tau_{ef}$ was 7.0 ms. In some cases (41%, n = 7/17) $\tau_{ef}$ was longer than the corresponding $eRT$, which suggests two things. First, the reset potentials were closer to the steady state mean $V_m$ than the natural fluctuations of $V_m$ around the mean, so the decay from reset back to the mean was faster than $\tau_{ef}$. Second, the $eAHP$ was close to zero or no more than a couple of milliseconds. This value of $eAHP$ is a dramatic decrease from the 200 ms AHP duration previously reported during quiescence [4,24]. Further, not only did the population spread in $eRT$ across cells correlate with the population spread in $SIT$ ($R^2 = 0.81$), the $eRT$ and the $eSIT$ both had significant correlation with $\tau_{ef}$ (Figure 5B) ($R_{SIT}^2 = 0.60$, $p = 0.0007$, $R_{eRT}^2 = 0.47$, $p = 0.005$, when ignoring the two outliers, cell 7 and 16). Since $eRT$ is dependent on both $\tau_{ef}$ and $eAHP$, while $eSIT$ is only dependent on $\tau_{ef}$ these strong correlations also indicate that the contribution of $eAHP$ to $eRT$ must be minor in most cells.

Hence, the population spread in $eRT$, $eSIT$, and $\tau_{ef}$ probably reflected different levels of synaptic intensity in different cells. Since the $eRT$ is the major contribution to the refractory period, we expected most of the inter-spike intervals (ISI) to be longer than or equal to the $eRT$. If this is the case, the data points in a plot of the shortest ISI versus $eRT$ in each cell should fall at the 45°-line.
Figure 4. Temporal characteristics of sub-threshold V_m-fluctuation in motoneurons during quiescence and in the on- and off-cycle during scratching. (A) Hip-flexor nerve recording during scratch. (B) Concurrent V_m in MN, spikes avoided with \(-2.5\) nA hyperpolarizing current. Shaded regions mark the selected area of on- and off-cycle illustration below. Sample trace of V_m in quiescence (C) (note time-course of spontaneous synaptic potentials) and in the on-cycle (D) and off-cycle (E). D and E are from the shaded boxes in B. (F) The auto-correlation sequence of each sample trace. Blue is from quiescent trace (C), gray is from the off-cycle trace (E), and red is the on-cycle trace (D). The effective time constant of each trace is obtained by fitting an exponential decay function (broken lines) to the initial 3 ms (until the vertical gray line). The time constants are \(t_{\text{eff}} = 2.8\) ms (on-cycle activity), \(t_{\text{eff}} = 5.2\) ms (off-cycle activity) and \(t_{\text{eff}} = 27.0\) ms (quiescence). C–E are on the same time scale.

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or below. Indeed, all points were within their error bars or below (Figure 5C). In this way, the inverse of the eRT represents an upper bound for the spike frequency during network activity.

Absence of spike frequency adaptation

The AHP contributes to spike frequency adaptation (SFA) in motoneurons at rest [4,24]. A classical example of SFA in absence of synaptic input is during bursting induced by N-Methyl-D-Aspartic acid (NMDA). We therefore performed a heuristic comparison between NMDA-bursting in MNs in slices with the bursting induced by synaptic input in functional networks (cf. Figures 6A and 6B). During each burst in the case of NMDA-bursting (Figure 6Aa), spike frequency peaked at onset and adapted to a lower level at the end of the burst (Figure 6Ab). The histogram of interspike-intervals confirmed similarity in firing rate distribution between NMDA-bursting and network bursting [28] (cf. Figure 6Bc and 6Ac). ISIN plotted against ISIN +1 displayed a ring-like pattern (Figure 6Ad), as expected from adapting spike trains in each successive burst [29].

The firing pattern in MNs during scratching was qualitatively different (Figure 6Bb). Though the distribution of ISI times resembled the NMDA-bursting (cf. Figure 6Bc and 6Ac), spike timing was irregular (Figure 6Bb) and points scattered nearly symmetrically around the 45°-line in the return map (51%/49%, SE = 6%) (Figure 6Bd). This shows that spike frequency acceleration was as prevalent as spike frequency adaptation during bursts [29] and therefore that the AHP and other intrinsic mechanisms for adaptation or acceleration in spike frequency did not have a detectable influence on the firing pattern during network activity. Furthermore, the correlation between ISI_N and ISI_{N+1} was marginal (R^2 = 0.03) (Figure 6Bc), which shows that spikes were driven by a stochastic process rather than by deterministic intrinsic properties. In all 17 motoneurons SFA was insignificant during depolarizing waves, i.e. the numbers above the line were not significantly higher than 50% (Figure 7). Thus, we conclude that the mechanisms causing SFA such as AHP accumulation were not pervasive enough to overcome the increase in synaptic conductance during motor activity.

Discussion

Spike timing, the principal output of neurons, is determined by interacting synaptic and intrinsic ionic conductances. Recent decades have provided a wealth of information about the intrinsic response properties and their proposed roles in specific cell types in many parts of the nervous system [5,30,31]. Properties of the AHP in particular have been linked to classification of fast and slow
Figure 6. Presence and absence of spike frequency adaptation (SFA) during bursting. (Aa) SFA in intracellular recording from motoneuron in slice during NMDA induced bursting. (Ab) Single burst highlighted in (Aa) show gradual increase in ISI. (Ac) Histogram of ISI. The mean ISI is 55 ms (arrow). (Ad) Plot of ISI$_n$ against ISI$_{n+1}$ shows significantly greater proportion of points above than below the ISI$_n$ = ISI$_{n+1}$ line (83.2% above, total N = 239), which is evidence of spike frequency adaptation. In addition, ISIs are correlated with their neighbors (correlation coefficient = 0.41), as expected when ISI are influenced by AHP conductance and the burst pattern is reproduced after 10 spikes (Ae). Gray area represents the 5% confidence limit $\pm 1.96/\sqrt{N}$. (Ba) Recording from a MN in a functional spinal network during rhythmic motor activity. (Bb) A highlighted cycle from (Ba) shows irregular spike times and no SFA. (Bc) Histogram of ISI. The mean ISI is 34 ms (arrow). (Bd) A plot of ISI$_n$ against ISI$_{n+1}$ illustrates no discrepancy of points above and below the ISI$_n$ = ISI$_{n+1}$ line (51.3% above, total N = 362), which demonstrates absence of SFA. Furthermore, there is only a marginal correlation of ISI with neighbors (correlation coefficient = 0.18), as expected with negligible AHP conductance (Be). Gray area represents the 5% confidence limit $\pm 1.96/\sqrt{N}$. Tick marks to the left represent $\sim$80 mV (A) and $\sim$60 mV (B). Inter-burst-intervals are omitted in analysis.

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spiking neurons with clear functional connotations [5,32]. The present study reports up to ten-fold decrease in the membrane time constant (Figure 4) and severe reduction in the AHP during functional activity (Figure 3 and 5) which emphasize that assertions about the functional significance of intrinsic response properties must be validated in neurons embedded in an active network. In our experimental paradigm, the role of the AHP in spike timing is essentially eliminated and replaced by rapid temporal integration in the high conductance state provided by intense synaptic input (Figures 3, 5, 6). We therefore propose that motor systems perform rapidly changing movements by letting intense synaptic input transiently supplant slow intrinsic properties when necessary.

Coincidence detection

This transition has previously been referred to as a transition from temporal integration to coincidence detection in sensory perception [33]. Konig and colleagues defined a coincidence detector as a neuron in which the integration time for synaptic potentials is short compared with the average ISI, and a temporal integrator if the reverse is true. Adopting this definition, the motoneurons in the present study were all coincidence detectors during motor activity, since their eSIT and eRT on average are an order of magnitude shorter than their average ISI (cf. Figure 5A and 6Bc, and see Figure S5). The functional effect and benefit of MNs working as coincidence detectors could be to minimize aberrant firing due to the background barrage of noisy synaptic input [28]. The potentially undesirable consequence of having such a coincidence detection scheme is irregular spike patterns, which we indeed did observe (Figure 6). However, irregular patterns are not likely to be a problem, since muscle fibers are slow integrators, and therefore the exact temporal structure of the MN firing, whether irregular and yet fast coincidence detectors or slow regular temporal integrators, is unimportant for securing a smooth contraction.

Origins of spike pattern

The network mechanisms underlying the phasic spike activity in motoneurons during rhythmic motor behaviors are unknown. It has been hypothesized that certain intrinsic properties are crucial mediators of bursting rhythms in spinal networks [3,5,34]. It is also widely accepted that the after-hyperpolarization in MN secures repetitive firing at low rates appropriate for regulation of muscle contraction [3,4]. The high conductance state, however, compromises the ability to fire repetitively during steady depolarization [11]. This is probably a direct consequence of the reduced slow AHP since the same effect is observed when the AHP is reduced by blocking the underlying ionic current pharmacologically rather than by shunting [35]. Nevertheless, MNs still fired in a broad range of rates in the high conductance state while driven by rapid fluctuations in membrane potential during motor activity (Figure 6B) [13]. Further evidence for fluctuation driven irregular firing is the fact that the shortest ISIs in the active network are longer than the eRT (and thus eAHP) (Figure 6C). This is in contrast to the regular firing and frequency adaptation in motoneurons in slices during NMDA induced bursting (Figure 6A). The reduced AHP amplitude, increased spike time variability and the absence of adaptation during locomotor network activity have also been observed in MNs in the decerebrate cat [36]. In the light of our findings it will be interesting to investigate if the underlying mechanism is modulation of intrinsic properties [1,24,37] or parallel increase in excitatory and inhibitory synaptic activity [13,38].

In conclusion, the rhythm-generation could have two origins: a pattern generating subset of neurons elsewhere in the network in which intrinsic response properties are protected from shunting by
intense synaptic bombardment or alternatively, the motor rhythm can be an emergent distributed network phenomenon as suggested for respiratory rhythms [39,40], i.e. the network bursting hypothesis. Decisive experimental tests of these hypotheses are still missing.

Caveats of constant current protocols

The role of AHPs in repetitive firing in MNs induced by depolarizing current through the recording electrode has been thoroughly investigated [4,5,24]. Based on the linear summation of synaptic potentials under certain experimental conditions [41,42] this approach takes injected current as a simplifying representation of synaptic input. However, important aspects of synaptic input are overlooked in this approach. First, synaptic variability adds fluctuations as a second moment to \( V_m \) [22]. Synaptic \( V_m \)-fluctuations may have important computational roles [43–45]. Secondly, the effect of surge in conductance from synaptic input is unaccounted for in the constant current protocol (Figure 1). Therefore the functional role of the AHP and other intrinsic response properties should be assessed either in functional networks with real synaptic input, as in the present study, or by dynamic clamp analysis in neurons at rest [7,8,46] if a valid estimate of the temporal structure of the functional synaptic conductances is available.

Balanced state in the spinal cord

Though anatomical evidence suggests an approximate balance between inhibitory and excitatory contacts in cat motoneurones [4], the circuit substrate for variations in the balance of inhibition and excitation during motor activity and its prevalence in other spinal networks is yet unresolved. High-conductance states produced by parallel increase in inhibitory and excitatory synaptic activity is a common occurrence in other functional networks [6,38,47–51]. In the neocortex, the link between excitation and inhibition is provided by feed-forward and recurrent pathways [48,49,52–55]. In the spinal cord, recurrent inhibition is unlikely to contribute significantly to the high conductance state in MNs since recurrent collaterals are scarce in the turtle [56,57] and Renshaw inhibition has not been documented. Feed-forward inhibition provided by Ia inhibitory interneurons can produce synaptic conductance that clearly reduces intrinsic response properties in cat MNs in vivo [58]. However, reciprocal inhibition contributes insignificantly to the high conductance state in MNs during breathing [38] and scratching [13]. We propose that inhibitory input to MN is mostly due to local feedforward inhibitory connections, but it remains to be seen if balanced spinal motor networks resemble architectures of the much better investigated networks in other parts of the nervous system [6,38,48,51,55].

Materials and Methods

All the experiments were performed in an integrated spinal cord-carapace preparation from the adult turtle except the heuristic control experiment of NMDA-induced spike frequency adaptation in figure 6A, which was performed in a transverse slice from adult turtle. In the integrated preparation the spinal cord remains in the spinal canal with the tactile sensory nerves from the carapace intact. The motor nerves are carefully transected to avoid muscle movements and dissected out for electro-neurogram recordings. The scratch reflex was activated by mechanical somato-sensory stimulation of selected regions on the carapace (see below) as described [11,20].

Integrated preparation

Red-eared turtles (Trachemys scripta elegans) were placed on crushed ice for 2 hrs to ensure hypothermic anesthesia. Animals were killed by decapitation and blood substituted by perfusion with a Ringer solution containing (mM): 120 NaCl; 5 KCl; 15 NaHCO3; 2 MgCl2; 3 CaCl2; and 20 glucose, saturated with 90% O2 and 2% CO2 to obtain pH 7.6. The carapace containing the D4-D10 spinal cord segments was isolated by transverse cuts and removed from the animals, similar to studies published elsewhere [19,20]. The surgical procedures complied with Danish legislation and were approved by the controlling body under the Ministry of Justice.

Slice preparation

One mm thick slices of the turtle spinal cord were placed in a chamber for intracellular recording and submerged in and perfused with oxygenated Ringer solution. The pharmacological agent N-methyl-D-aspartate (NMDA) was added to the ringer medium to induce bursting activity (10 µM).

Recordings

Intracellular recordings in current-clamp mode were performed with an Axoclamp-2A amplifier (Axon Instruments, Union City, CA). Glass pipettes (part no. 30-0066, Harvard Apparatus, UK) were pulled with a electrode puller (model P-87, Sutter instrument co., USA) and filled with a mixture of 0.9 M potassium acetate and 0.1 M KCl. Intracellular recordings were obtained from neurons in segment D10. Recordings were accepted if neurons had a stable membrane potential more negative than –50 mV. Data were sampled at 20 kHz with a 12-bit analog-to-digital converter (Digidata 1200, Axon Instruments, Union City, CA), displayed by means of Axoscope and Clampex software (Axon Instruments, Union City, CA), and stored on a hard disk for later analysis. Hip flexor nerve activity was recorded with a differential amplifier Iso-DAM8 (WP) using a suction pipette. The bandwidth was 100 Hz–1 kHz.

Activation of network

Mechanical stimulation was performed with the fire polished tip of a bent glass rod mounted to the membrane of a loudspeaker in the cutaneous region known to elicit “pocket scratch” [12] which results in a broad activation of cells [59]. The duration, frequency, and amplitude of the stimulus were controlled with a function generator (Figure 2A). This tactile stimulus induced the scratch-like network activity, which was monitored by the suction electrode nerve recordings from the Hip-flexor nerve (Figure 2B).

Model data

The data used to illustrate the difference between synaptic-current and synaptic-conductance based fluctuating inputs (Figure 1) was based on a one-compartment model simulation [22] supplemented with Ca2+-conductance and a Ca2+-activated K+-conductance. The synaptic noise was modeled as white current noise in the current-based regime with the heuristic expression

\[
\frac{dV_m}{dt} = \frac{G_{leak}(E_{leak} - V_m) + G_{AHP}(E_K - V_m) + I_{syn}}{C}
\]

where the fast conductances of the action potential and Ca2+-conductance were not shown here for simplicity (for complete description see Data and Methods S1) and the total membrane conductance was

\[
G_{total} = G_{leak} + G_{AHP}
\]

which contains no synaptic component. In this scheme, the synaptic input was represented as a current, \( I_{syn} \). The membrane capacitance is \( C \), and \( G_{leak} \). \( E_{leak} \), \( G_{AHP} \), \( E_K \) are conductance and reversal potential of leak and AHP, respectively.
In a more realistic regime, the high intensity synaptic input was modeled as a conductance [22]. Here, the AHP was reduced as a consequence of the increase in total input conductance (Figure 1). The conductance-based regime applies when the synaptic conductance (G_syn) is so large that it can no longer be considered small compared with the G_total [6]. Heuristically expressed similar to the synaptic current model

$$\frac{dV_m}{dt} = \frac{G_{\text{leak}}}{G_{\text{total}}} (V_m - E_{\text{leak}}) + \frac{G_{\text{AHP}}}{G_{\text{total}}} (V_m - E_{K}) + \frac{G_{\text{Syn}}}{G_{\text{total}}} (V_m - E_{\text{syn}})$$

where $E_{\text{syn}}$ is the weighted reversal potential of excitatory and inhibitory synaptic reversal potentials and $G_{\text{syn}}$ is the sum of both conductances. $G_{\text{syn}}$ is competing with $G_{\text{AHP}}$ in controlling interspike intervals, and if it is large enough it can render $G_{\text{AHP}}$ insignificant at steady state:

$$V_m = \frac{G_{\text{AHP}}}{G_{\text{total}}} E_{K} + \frac{G_{\text{leak}}}{G_{\text{total}}} E_{\text{leak}} + \frac{G_{\text{Syn}}}{G_{\text{total}}} E_{\text{syn}} \approx \frac{G_{\text{leak}}}{G_{\text{total}}} E_{\text{leak}}$$

when $G_{\text{Syn}} \gg G_{\text{AHP}}$. The action potentials were largely unaffected by the conductance increase, since fast Na/K conductances of the action potential were largely unaffected by the conductance increase, since fast Na/K conductances of the action potential was more than 12 nA, $\sigma_{\text{syn}} = 3$ nA, OU-simulated with time-constant = 1 ms and D = 0.0005) or conductance noise we verified the theoretical importance of synaptic conductance (Figure 1). The conductance noise was a mixture between inhibition and excitation balanced at $-60$ mV (see Data and Method S1 for details).

**Evolution of $V_m$-distribution**

The variable constituting the estimation of effective eSIT and eRT was the membrane potential ($V_m$). This variable was stochastic and a sample measurement drawn from an underlying probability distribution function (PV), which we assumed had the same statistics in all interspike intervals. The probability distribution depends on time after occurrence of spike and this dependence was a manifestation of intrinsic current generators like SK-channels. Because of the large synaptic fluctuations, it was necessary to look at the distribution PV instead of just isolated instances of $V_m$. These fluctuations were assumed uncorrelated from trial to trial, so we could estimate $P_v$ by superimposing spikes.

The key assumption is, if the distribution at some given point in time, $P_v(t_1)$, is different from the distribution at a later point in time, $P_v(t_2)$, then there has been a change in the intrinsic current generation (cf. Figure 3A and B). We selected $P_v$ at one point in time ($t = t_{\text{template}}$) as a template distribution, which all distributions $P_v(t \neq t_{\text{template}})$ were compared with. The $P_v(t_{\text{template}})$ was chosen more than 10 ms before the spike, since this region constitute a background $V_m$ and was compared via the Kolmogorov-Smirnov-2 sample test (KS-test) [60] expressed formally as:

$$F(V, t) = \frac{\text{Number of } V_m(t) \leq V}{N}$$

$$D(t) = \sup \left| F(V, t) - F(V, t_{\text{template}}) \right|$$

$F(V, t)$ is the empirical cumulative probability distribution function of $V_m$, N is the total number of traces (and spikes) used to estimate the distribution at time t (Figure 3D). The maximal difference between the cumulatives, $D(t)$, was the measure for rejecting or accepting the null hypothesis. If the difference was larger than a critical value, then we rejected the null hypothesis that $V_m(0)$ and $V_m(t_{\text{template}})$ were drawn from the same distribution. The binary test outcome was plotted (Figure 3A and B) where 1 represented rejection (gray area) and 0 represented no rejection of null hypothesis, at a 5% confidence limit. The p-value of the test was plotted below.

**Effective Recovery time**

The first point in time after the spike, where the KS-test was zero (i.e., no rejection of hypothesis of same distribution) was where we defined the AHP conductance and other transient intrinsic current generators no longer had a significant impact on the $V_m$ and the passive diffusive spread had reach steady state. We dubbed this period *effective recovery time* (eRT, arrow in Figure 3B) in analogy to the effective membrane resistance and effective membrane time constant [14,16,21,22]. This recovery time told us how long time after the spike had occurred that there was still a memory of the spike in $V_m$ (Figure 3B).

**Effective synaptic integration time**

Similar to eRT, we could ask how long time prior to the spike, that $P_v$ was different from the template distribution. This point represented a net depolarization caused either by reduced inhibition or increased excitation. We named this period *effective synaptic integration time* (eSIT) indicating the time prior to the spike where its occurrence can be predicted (arrow in Figure 3B).

**Location of template**

Obviously, the choice of template distribution is important. The template is always chosen prior to the spike. The earlier before the spike we choose the template, the more independent it is. However, there is a trade off, since the inter-spike interval has to be longer than the window between the template distribution and the spike. As a result, the larger the window is, the fewer spikes in a finite dataset will participate in the distribution (Figure 3D). Fortunately, the estimation of both eRT and eSIT is largely independent on the window size (Figure 3C). We chose to average the values of eRT and eSIT from templates 20 to 40 ms prior the spike, since these locations gave nearly constant values (Figure 3C).

**Critique of method**

The above described statistical testing of the evolution of $P_v$ only accounts for changes that are locked to the occurrence of a single action potential, such as the AHP. Accumulative events that build up over several spikes as e.g. spike frequency adaptation or plateau potentials are not easily accounted for using this statistics. One way to test for slow changes would be to divide the spikes according into several different groups depending on their position in the epoch. These groups could then be compared to evaluate if the distributions have changed. However, we decided this was outside the scope of the present study and it was not necessary since the test of spike frequency adaptation (Figure 6) came out negative.

**Inter-spike interval analysis**

The inter-spike intervals were extracted from the intracellular recording during scratch episodes and processed. The auto-correlations were calculated as the normalized covariance function [26,61] between ISIn and ISIn+1, ISIn+2, ISIn+3 etc (Figure 6d). The test for spike frequency adaptation (Figure 6c) was performed...
assuming statistical independence of observation of \((ISIN, ISIN_{\pm 1})\)-pairs [29] and thus a binomial distribution with chance of 50% above and 50% below the line \(ISIN = ISIN_{\pm 1}\). If the number of points above was within a standard deviation of

\[
\sigma = \sqrt{Kp(1-p)} = \sqrt{K \cdot 0.25} \quad \text{SE} = \sqrt{K \cdot 0.25} / K = \frac{0.25}{\sqrt{K}}.
\]

of the 50%-line, as expected from an even binomial distribution [61], there was no significant spike frequency adaptation. \(K\) is the total number of points and \(p\) is the probability of a point being above the line, when assuming no correlation.

Data processing

All analysis was performed in Matlab (version 7.3, Mathworks). The data was converted from Axoclamp format to matlab and the spikes were identified and superimposed (Figure 3) to study the \(V_m\) statistics before and after the spike as described above. KS-testing of the \(V_m\) distributions was done with the matlab procedure “kstest2.m”. The custom made procedures for calculating eRT and eSIT has been uploaded to mathworks code sharing web site (http://www.mathworks.com/matlabcentral/) with the name “eRT.m” for the interested reader. The remaining matlab code is available on request.

References

1. Miles GB, Hartley R, Todd AJ, Brownstone RM (2007) Spinal cholinergic interneurones regulate the excitability of motoneurones during locomotion. PNAS 104: 2448–2453.
2. Rekling JC, Funk GD, Bayliss DA, Dong XW, Feldman JL (2000) Synaptic control of motoneuronal excitability. Physiological reviews 80: 767–852.
3. Grillner S (2006) Biological pattern generation: The cellular and computational logic of networks in motion. Nature 52: 751–766.
4. Kornell D (2006) The motoneuron and its muscle fibres. Oxford.
5. Toledo-Rodriguez M, Ei Manira A, Wallen P, Svirskis G, Hounsgaard J (2005) Cellular signalling properties in microcircuits. TINS 28: 534–540.
6. Destexhe A, Rudolph M, Pare D (2003) The high-conductance state of neocortical neurons in vivo. Nature Rev Neurosci 6: 739–751.
7. Wolfart J, Dehuy D, Masson GL, Destexhe A, Bal T (2005) Synaptic background activity controls spike transfer from thalamus to cortex. Nature Neuroscience 8: 1760–1767.
8. Fernandez FR, White JA (2008) Artificial Synaptic Conductions Reduce Subthreshold Oscillations and Periodic Firing in Stereotyped Cells of the Entorhinal Cortex. J Neuroscience 28: 3790–3803.
9. Borg-Graham LJ, Monier C, Fregnat Y (1996) Voltage-clamp measurement of visually-evoked conductions with whole-cell patch recordings in primary visual cortex. J Physiology (Paris) 90: 185–188.
10. Borg-Graham LJ, Monier C, Y. F (1998) Visual input evokes transient and strong shunting in inhibition in cortical neurones. Nature 393: 369–373.
11. Alaburda A, Russo R, MacAulay N, Hounsgaard J (2005) “Periodic high-conductance states in spinal neurones during scratch-like network activity in adult turtles”. J Neurosci 25: 6316–6321.
12. Robertson GA, Stein PSG (1968) Synaptic control of hindlimb motoneurones during three forms of fictive scratch reflex in the turtle. J Physiology 404: 101–129.
13. Berg RW, Alaburda A, Hounsgaard J (2007) Balanced inhibition and excitation drive spike activity in spinal half-centers. Science 315: 390–393.
14. Barrett JN (1975) “Motoneuron dendrites: role in synaptic integration.” Federation proceedings 34: 1396–1407.
15. Destexhe A, Rudolph M, Fellous J-M, Sejnowski TJ (2001) Fluctuating synaptic conduances recreate in vivo-like activity in neocortical neurons. Neuroscience 107: 13–24.
16. Bernardner OJDR, Martin KAC, Koch C (1991) Synaptic background activity influences spatiotemporal integration in single pyramidal cells. PNAS 88.
17. Shefchik SJ, Jordan LM (1985) Motoneuron input-resistance changes during fictive locomotion produced by stimulation of the mesencephalic locomotor region. J Neurophysiol 54: 1101–1108.
18. Perreault MC (2002) Motoneurons have different membrane resistance during fictive scratching and weight support. J Neurosci 22: 8259–8265.
19. Keifer J, Stein PSG (1983) Invitro motor program for the rostral scratch reflex generated by the turtle spinal-cord. Brain research 266: 148–151.
20. Alaburda AJH (2003) Metabotropic modulation of motoneurones by scratch-like spinal network activity. J Neurosci 24: 8923–8929.
21. Rapp M, Yarom Y, Seges I (1992) The impact of parallel fiber background activity on the cable properties of cerebellar purkinje cells. Neural Computation 4: 310–333.
22. Tiesinga PHE, Jose JV, Sejnowski TJ (2000) Comparison of current-driven and conductance-driven neocortical models with Hodgkin-Huxley voltage-gated channels. Physical review E 62: 8413–8419.
23. Destexhe A, Pare D (1999) Impact of network activity on the integrative properties of neocortical pyramidal neurones in vivo. J Neurophysiol 81: 1531–1547.
24. Brownstone RM (2006) Beginning of the end: repetitive firing properties in the final common pathway. Progress in Neurobiology 78: 156–172.
25. Guillamon A, McLaughlin DW, Rinzel D (2006) Estimation of synaptic conductances. J Physiology Paris 100: 31–42.
26. Tuckwell HC. Introduction to theoretical neurobiology: Nonlinear and stochastic theories: Cambridge University Press.
27. Ditlevsen S, Lansky P (2003) Estimation of the input parameters in the Ornstein-Uhlenbeck neuronal model. Physical review E 71: 011907.
28. Mazzoni A, Broccard FD, Garcia-Perez E, Beniﬁzi P, Ruoaro ME, et al. (2007) On the dynamics of the spontaneous activity in neuronal networks. PLoS one 2: e189.
29. Rodieck RW, Kang NY-S, Gerstein GL (1962) Some quantitative methods for the study of spontaneous activity of single neurones. Biophysical. J. 2: 351–368.
30. Llinás RR (1988) The intrinsic electrophysiological properties of mammalian motoneurones. J Physiol 404: 115–137.
31. Konig P, Engel AK, Singer W (1996) Integrator or coincidence detector? The role of the cortical neuron revisited. TINS 19: 130–137.
32. Bartos M, Vida I, Jonas P (2007) “Synaptic mechanisms of synchronized gamma oscillations in inhibitory interneuron networks”. Nat Rev Neurosci 8: 45–56.
33. Hounsgaard J, Kiehn O, Mintz I (1988) Adapting motoneurones for motor behavior. Progress in Brain Research 23: 57–67.
34. Alaburda A, Vinh DM, Hounsgaard J (1999) Role of the spinal network activity in the generation of fictive locomotion. J Physiol 510: 2448–2453.
35. Hounsgaard J, Kiehn O, Mintz I (1988) Response properties of motoneurones in vivo. J Physiol 398: 573–589.
36. Brownstone RM, Jordan LM, Kriellaars DJ, Noga BR, Shefchik SJ (1992) On the regulation of repetitive firing in hindlimb motoneurones during fictive locomotion in the cat. Exp Brain Res 90: 441–453.
37. Schmidt BJ (1994) Afterhyperpolarization modulation in lumbar motoneurons during locomotor-like rhythmic activity in the neonatal rat spinal cord in vitro. Exp Brain Res 99: 214–222.

38. Parkis MA, Dong X-W, Feldman JL, Funk GD (1999) Concurrent inhibition and excitation of phrenic motoneurons during inspiration: Phase-specific control of excitation. J Neuroscience 19: 2368–2380.

39. Del Negro CA, Morgado-Valle C, Feldman JL (2002) Respiratory rhythm: an emergent network property? Neuron 34: 821–830.

40. Feldman JL, Del Negro CA (2006) Looking for inspiration: New perspectives on respiratory rhythm”. Nat Rev Neurosci 7: 232–241.

41. Granit R, Kernels D, Lamarre Y (1966) Algebraic summation in synaptic activation of motoneurones firing in the “primary range” to injected currents. J Physiol 187: 379–399.

42. Powers RK, Binder MD (2000) Summation of effective synaptic currents and firing rate modulation in cat spinal motoneurons. J Neurophysiol 83: 483–500.

43. Chance FS, Abbott LF, Reyes AD (2002) Gain modulation from background synaptic input. Neuron 35: 773–782.

44. Shadlen MN, Newsome WT (1996) The variable discharge of cortical neurons: Implications for connectivity, computation, and information coding. J Neurosci 16: 3870–3896.

45. Stein RB, Gersen ER, Jones KE (2005) Neuronal variability: Noise or part of the signal? Nat Rev Neurosci 23: 4985–4995.

46. Prinz AA, Abbott LF, Marder E (2004) The dynamical clamp comes of age. TINS 27: 218–224.

47. Csicsvari J, Hirase H, Csukro A, Mamiya A, Buzsaki G (1999) Oscillatory coupling of hippocampal pyramidal cells and interneurons in the behaving rat. J Neurosci 19: 274–287.

48. Haider B, Duque A, Hasenstaub AR, McCormick DA (2006) Neocortical network activity in vivo is generated through a dynamical balance of excitation and inhibition. J Neurosci 26: 4535–4545.

49. Wehr M, Zador AM (2003) Balanced inhibition underlies tuning and sharpens spike timing in auditory. Nature 426: 442–446.

50. Baca SM, Marin-Burgin A, Wagenaar DA, Kristan WB Jr (2008) Widespread inhibition proportional to excitation controls the gain of a leech behavioral circuit. Neuron 57: 276–289.

51. Monier C, Fournier J, Fregna Y (2008) In vitro and in vivo measures of evoked excitatory and inhibitory conductance dynamics in sensory cortices. J Neurosci 169: 323–365.

52. Kafiper C, Glickfeld LL, Atallah BV, Scanziani M (2007) Supralinear increase of recurrent inhibition during sparse activity in the somatosensory cortex. Nature Neuroscience 10: 743–753.

53. Silberberg G, Markram H (2007) Disynaptic inhibition between neocortical pyramidal cells mediated by Martinotti cells. Neuron 53: 735–746.

54. Monier C, Chavane F, Bandet P, Graham LJ, Fregna Y (2005) Orientation and direction selectivity of synaptic inputs in visual cortical neurons: a diversity of combinations produces spike tuning. Neuron 37: 663–680.

55. Marino J, Schummers J, Lyon DC, Schwabe L, Beck O, et al. (2005) Invariant computations in local cortical networks with balanced excitation and inhibition. Nature Neuroscience 8: 194–201.

56. Ruigrok TJ, Crowe A, Donkelaar HJ (1984) Morphology of lumbar motoneurons innervating hindlimb muscles in the turtle Pseudemys scripta elegans: an intracellular horseradish peroxidase study. J Comp Neurosci 230: 413–425.

57. Chmykhova NM, Aftanina VO, Karamian OA, Kozhanov VM, Vesselkin NP, et al. (2003) Comparative study of spinal motoneuron axon collaterals. Brain Research Bulletin 66: 381–386.

58. Hyngstrom AS, Johnson MD, Miller JF, Heckman CJ (2007) Intrinsic electrical properties of spinal motoneurons vary with joint angle. Nature Neuroscience 10: 363–369.

59. Berkowitz A (2001) “Broadly tuned spinal neurons for each form of fictive scratching in spinal turtles”. J Neurophysiol 86: 1017–1026.

60. Press WH, Teukolsky SA, Vetterling WT, Flannery BP (1992) Numerical recipes in fortran-The art of scientific computing. 2nd edition Cambridge University Press.

61. Taylor JR (1982) An introduction to error analysis-The study of uncertainties in physical measurements. California: University Science books.