Synaptic Interactions Between Forelimb-Related Motor Cortex Neurons in Behaving Primates

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Smith WS, Fetz EE. Synaptic interactions between forelimb-related motor cortex neurons in behaving primates. J Neurophysiol 102: 1026–1039, 2009. First published May 13, 2009; doi:10.1152/jn.91051.2008. We investigated the synaptic interactions between neighboring motor cortex cells in monkeys generating isometric ramp-and-hold torques about the wrist. For pairs of cortical cells the response patterns were determined in response-aligned averages and their synaptic interactions were identified by cross-correlation histograms. Cross-correlograms were compiled for 215 cell pairs and 84 (39%) showed significant features. The most frequently found feature (65/84 = 77%) was a central peak, straddling the origin and representing a source of common synaptic input to both cells. One third of these also had superimposed lagged peaks, indicative of a serial excitatory connection. Pure lagged peaks and lagged troughs, indicative of serial excitatory or inhibitory linkages, respectively, both occurred in 5% of the correlograms with features. A central trough appeared in 13% of the correlograms. The magnitude of the synaptic linkage was measured as the normalized area of the correlogram feature. Plotting the strength of normalized interaction against response similarity during alternating wrist torques revealed a positive relationship for the correlogram area of the cell pairs. A linear fit yielded a positive slope: the pairs with excitatory interactions tended to covary more often than countervary. This linear fit had a positive offset, reflecting a tendency for both covarying and countervarying cells to have excitatory common input. Plotting the cortical location of the cell pairs showed that the strongest interactions occurred between cells separated by <400 microns. The correlational linkages between cells of different cortical layers showed a large proportion of common input to cells in layer V.

INTRODUCTION

The operations of cortical circuits are mediated by the synaptic interactions between cortical neurons, which depend on the strength and distribution of the functional connections between cells. Our understanding of intrinsic cortical circuitry has been significantly advanced by in vitro studies of synaptic interactions between morphologically characterized cells (Douglas and Martin 2004; Markram 1997; Markram et al. 1997; Thomson and Lamy 2007; Thomson et al. 2002). Yet, surprisingly little is known about normal interactions between functionally identified cortical neurons in vivo. Such interactions are sometimes inferred indirectly from response properties (Merchant et al. 2008). Synaptic interactions can be documented directly by cross-correlating the extracellular spike activity of the cells (Moore et al. 1970; Perkel et al. 1967) and by spike-triggered averaging of intracellularly recorded membrane potentials (Matsumura et al. 1996). Since intracellular recordings of synaptic potentials are difficult to obtain in behaving animals, cross-correlation is the more practical measure in the behavioral context required to define the cells’ normal functional properties. By measuring the synaptic interactions between cells as well as their task-related activity we can determine the extent to which the connections are functionally consistent with their response properties and to infer the mode of operations of cortical circuits. As reviewed elsewhere (Fetz et al. 1991) this strategy has been successfully applied in various cortical regions, including motor cortex.

Previous studies of precentral cells using correlation analysis have revealed evidence of positive synchronization between cells with similar response characteristics and negative correlation between cells that respond reciprocally (Allum et al. 1982; Georgopoulos et al. 1993; Jackson et al. 2003; Murphy et al. 1985), supporting the hypothesis that synaptic connections between precentral neurons shape the neurons’ discharge. Anatomical connections producing synchrony, such as recurrent synapses of pyramidal cells and radial afferent fibers, have also been suggested or identified (Ghosh and Porter 1988a,b; Kang et al. 1988; Stefanis and Jasper 1964), although the cell population they innervate has not been characterized.

This study documents the synaptic relationship between functionally identified precentral cell pairs and their response patterns during voluntary isometric wrist torques. In contrast to previous work, this study determined whether the interacting cells had axonal projections to spinal cord or correlational linkages with muscles. The study also provides the histological reconstruction of recording sites, elucidating the anatomical basis of synaptic interactions between cortical layers. The companion paper examines synaptic relations between identified corticomotoneuronal (CM) cells, which have correlational linkages with identified target muscles.

METHODS

Experiments were performed with two Macaca mulatta monkeys. The animals were housed, fed, and medically supervised at the Washington National Primate Research Center in conformity with U.S. Department of Health and Human Services guidelines for the care and use of laboratory animals. Monkeys were trained to perform isometric ramp-and-hold wrist torques in a task that has been used to characterize cells in cerebral cortex, red nucleus, and spinal cord (Cheney et al. 1991; Fetz et al. 2002). A 22-mm-diameter recording chamber was placed over the contralateral arm/hand precentral cortex and a chronically indwelling, concentric stimulating electrode placed in the ipsilateral medullary pyramid for antidromic activation of pyramidal tract (PT) neurons.
To facilitate the recording of cells within close proximity, we used a dual-electrode drive holder (Fig. 1). One electrode was advanced perpendicular to the chamber, whereas the other could be positioned at a 0–45° angle to the vertical electrode. The plane defined by the two electrodes could be translated mediolaterally and anteroposteriorly and the separation between both electrodes could be adjusted. The plane of the electrodes was maintained at a 30° angle to the anatomic sagittal plane, approximately perpendicular to the central sulcus. (This plane was chosen to maximize the possibility of observing serial inhibition produced by basket cells that send axons perpendicular to the long axis of the precentral gyrus.) Each glass-coated tungsten electrode was attached to a manually controlled hydraulic microdrive (Narashigi model 95A) and could be moved with 1-μm precision. After the electrodes penetrated the dura the chamber was filled with 10% sterile agar, which solidified and prevented displacement of the cortex by changes in intracranial pressure.

Each electrode signal was amplified by a band-pass amplifier (300 Hz to 5 kHz) and monitored by stereo headphones, with one channel for each signal. Once isolated, all cells were tested for an antidromic response to PT stimulation. The cell was classified as a PT neuron if it could be activated antidromically by a single stimulus through the medullary electrode and if this response could be collided with spontaneous, orthodromic spikes. Cells that did not meet both of these criteria were classified as non-PT (NPT) cells. As described in the companion paper (Smith and Fetz 2009), cells were identified as “CM cells” if spike-triggered averages (STAs) of rectified electromyographic (EMG) activity indicated a postspike facilitation of EMG.

At the end of each recording session, the electrodes were withdrawn to their initial extracranial position and the adaptor was removed from the monkey. The relative locations of the recording sites were reconstructed by repositioning the electrodes to the depths at which specific cells had been recorded in vivo and measuring the tip locations in reference to a 100-μm-resolution grid. Reconstruction accuracy was based on histological reconstruction of tracks in which electrolytic lesions were placed and further based on the measured electrode tip separation in cases when both electrodes recorded from the same cortical unit (10 instances). The accuracy of the electrode coordinates was estimated to be 135 ± 170 μm.

Signals from both electrodes, wrist torque, and EMG activity of forearm muscles were recorded on a frequency-modulated tape recorder for off-line analysis. Action potentials of cortical units were used to generate digital pulses using time–amplitude window discriminators (Model DIS-1; Bak Electronics); waveform separation was sometimes enhanced with an audio equalizer (Model 31–2009; Radio Shack). These discriminated spike trains were digitized with 250-μs resolution along with the analog wrist torque signal at 20-ms resolution. A unit was excluded from further analysis if the autocorrelogram of its spike train revealed counts within the refractory period (typically within 3 ms of the origin), indicating contamination from other cortical cells.

“Response average” histograms of unit activity aligned with the ramp portion of wrist torque were generated for all cells. To quantify the degree to which the firing rates of two units covaried during the task, we calculated a measure called “response similarity,” as illustrated in Fig. 2A. Each unit’s spike train was converted to a continuous waveform by summing all spikes that occurred within a moving 200-ms window, advanced every 20 ms. This sum was placed into a bin at the center of the 200-ms window. To remove offset, the mean of the entire trace was subtracted from each bin. A scatterplot of the two smoothed firing rates in corresponding bins was generated (top right). Response similarity was defined as the regression coefficient “r” for these points. In this example the points fell along a line with a positive slope, producing a positive r value (0.87). Units whose firing varied reciprocally had a negative linear regression and a negative r value. Scatterplots of units with uncorrelated firing generated a circular pattern and produced a “response similarity” near zero.

Correlograms between spike trains of cell pairs were constructed by the method of Perkel et al. (1967). Correlograms of 1-ms and 250-μs bins were calculated from raw spikes from cell pairs during performance of alternating wrist torques. Periods of inactivity, sleep, and restlessness were excluded from analysis.

The resulting correlograms were inspected by eye to identify sustained deviations from baseline. Baseline was defined as the first 50 bins of the 200-bin correlogram (Fig. 2B). Onset and offset of a feature were defined as the central bin of a three-bin average that exceeded the baseline mean (in the case of peaks) or that fell below this mean (in the case of troughs). All bins in the feature satisfied the three-bin average criterion. Single bins with deviant counts, which usually represented cross talk between electrodes, were excluded if the counts failed to distribute across adjacent bins when the bin resolution was quadrupled to 250 μs. The time between the onset and offset bins defined the feature duration. Area of the feature was defined as the total number of counts above baseline in bins between onset and offset, inclusively. The absolute area was tested for statistical significance using the Z statistic (Cope et al. 1987). Correlograms with |Z| ≥ 2.8 (P < 0.005) were accepted as significant.

For all correlograms the feature area was divided by the algebraic mean of the number of target and reference cell spikes that occurred during construction of the cross-correlogram. This measure is referred to as normalized area or AN. Thus AN is the feature area divided by (Np + Nr)/2, where Np is the number of reference cell triggers and Nr is the number of target cell triggers. AN represents the probability that any one spike, from either cell, is correlated above chance with the other cell. We also calculated the parameters α = AN/Nr and contribution = AN/Nr.
The relative locations of recorded units within cortical laminae and their separation distance were determined from histological reconstructions for every cell pair. The cortex of the first animal was reconstructed from cresyl-stained 60-μm sections. In the second animal the corticospinal neurons were retrogradely labeled. Two days before sacrifice, 2 mg of wheat-germ agglutinin conjugated horseradish peroxidase (WGA-HRP) was injected into the lateral lamniscus, intermediate horn, and lateral-ventral horn of the contralateral spinal segments C4–T1. Brain sections were processed using an ammonium heptahydrate molybdate tetramethyl benzidine procedure to stain retrogradely transported HRP (Olucha et al. 1985). Features of each cortical section were digitized using a microscope-optical imaging program, which included the cortical surface, location of layer V, gray–white junction, and the locations of HRP-stained neurons (shown as dots in Fig. 1). These digitized sections were then translated into coordinates of the chronic recording chamber, allowing the superposition of recording positions over the appropriate histologic area.

Since thalamocortical and corticocortical afferents arborize radially, irrespective of gyrus morphology, a useful measure of cell separation is the horizontal separation between the columns in which each cell resides. To measure this distance, the histologic section was optically projected over the computer-reconstructed electrode tracks in register with the digitized histologic section. Each cell of the pair was projected to layer V along radial fibers and a line integral along layer V between these projected points was calculated using 50-μm calipers. This distance is called “horizontal separation.”

The presumptive laminar locations of all cells were identified by superimposing electrode tracks over the appropriate histologic section. Recording sites of each cell were classified as being located within either laminae II–III, V, or VI. Laminae II and III were grouped together because discrimination of the cytoarchitectonic boundary between them was difficult with the stains used. The Nissl stain used in the histology of animal 1 allowed clear delineation of layer VI (below layer V and absence of Betz cells), layer V (presence of Betz cells), and layer II–III (above layer V). In animal 2, the neutral red counterstain made delineation of the superficial border of layer V difficult; when a cell was judged to be on the border of layer III–V, the cell was categorized as a layer V cell. Layer VI cells were easily identified in this monkey since the deep border of layer V was well defined by the population of labeled corticospinal cells.

RESULTS

Database and classification of correlogram features

While the monkeys performed alternating ramp-and-hold wrist torques, a total of 221 precentral cells were isolated in a total of 110 electrode tracks. Cross-correlograms were compiled for 215 cell pairs. Thirty-nine percent of these correlograms (n = 84) exhibited significant features. A feature was defined as a significant positive or negative deviation of the correlogram from baseline that was sustained for successful bins. Five types of correlogram features were observed. 1) Central peaks straddled the origin (the time of the trigger cell spike) and dropped off more or less symmetrically; i.e., the time of peak was usually within 1 ms of the origin and the rate of decay was similar in the positive and negative directions. In contrast, 2) lagged central peaks had peaks displaced from the origin and demonstrated appreciable asymmetry in decay for

FIG. 2.  A: calculation of response similarity (r), measuring the strength of covariation in firing rates of 2 precentral cells during active wrist torque generation. The digitized spike trains of 2 neurons (units 25-4 and 27-2) and wrist torque are shown for 10 s of data. The units’ firing rate is smoothed by summing the number of spikes within a moving 200-ms window as shown in the "smoothed" traces. Response similarity is calculated as the correlation coefficient between the 2 smoothed signals, shown in the scatterplot on the right (r = 0.87). Torque signal: flexion upward, extension downward. B: determination of onset, offset, and size of correlogram feature. Baseline was defined as the first 50 bins of the correlogram, onset, and offset as the middle bin of a 3-bin average that exceeded this baseline (for peaks) or fell below (for troughs). Area is the sum of counts above or below baseline between onset and offset: AN is the feature area divided by (NR + NT)2, where NR is the number of reference cell triggers and NT is the number of target cell triggers. Features with |Z| > 2.8 were accepted as significant (Cope et al. 1987). Numerical measures for this example are tabulated. α = AN/NT; contribution = AN/NT.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Mean N bins</th>
<th>Onset Offset Width (ms)</th>
<th>Baseline:</th>
<th>Peak:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base</td>
<td>41.5</td>
<td>-100 -50</td>
<td>51</td>
<td>51</td>
</tr>
<tr>
<td>Peak</td>
<td>50.1</td>
<td>-10 +11</td>
<td>22</td>
<td>22</td>
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<table>
<thead>
<tr>
<th>Area (cm²)</th>
<th>Alpha</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>191</td>
<td>0.111</td>
<td>0.118</td>
</tr>
<tr>
<td>4.94</td>
<td>0.105</td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>N_R</th>
<th>N_T</th>
<th>Z</th>
<th>Area</th>
<th>Alpha</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1616</td>
<td>1817</td>
<td>4.94</td>
<td>191</td>
<td>0.111</td>
<td>0.118</td>
</tr>
</tbody>
</table>

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positive and negative lag times. 3) Central troughs were sustained deviations below baseline that straddled the origin. 4) Lagged peaks were positive deviations above baseline with onsets after the correlogram origin. Finally, 5) lagged troughs were sustained negative deviations with onset after the reference spike trigger that subsequently returned to baseline.

All 215 correlograms were first analyzed by eye to detect the potential presence of any of these five features. Onset and offset of the feature and the area above (or below) baseline comprising the feature were subjected to statistical tests (Z statistic; see METHODS). In correlograms without overt features, a standard ±10-ms range was adopted for statistical testing. Features with \(|Z| \geq 2.8\) \((P < 0.005)\) were considered significant; this correlated well with those judged to be significant by eye.

Examples of cross-correlogram features

Figures 3–7 show examples of each of the five types of correlogram feature with the response averages of each cell comprising the pair and the cortical location of the cells. Central peaks represented the most frequent correlogram feature type, found in 44/84 significant correlograms. The example in Fig. 3 shows a correlogram with a central peak of 21-ms duration (horizontal bar below peak). These two PT cells were recorded in electrode tracks 25 and 27 (arrows, bottom left diagram) separated by 118 \(\mu\)m within layer V on the convexity of the gyrus. The presence of a central peak in the cross-correlogram can be interpreted as evidence for common synaptic input (Dickson and Gerstein 1974; Perkel et al. 1967), so a common afferent is shown connecting the two cortical cells (see DISCUSSION). The response averages (bottom right) indicate that the activity of both cells increased during extension torques and declined during flexion, in a parallel manner, yielding a response similarity of 0.89.

Lagged peaks were seen in isolation in 4/84 significant correlograms. Figure 4 illustrates a cell pair recorded simultaneously on the same electrode with a lagged correlogram peak. Both of these NPT cells were recorded deep in the bank of the precentral gyrus, below the HRP-labeled corticospinal population of layer V neurons (shown as dots). As in Fig. 3, the response profiles of the two units are very similar, differing mainly in absolute firing rate. The presence of a lagged peak with onset within 1.25 ms of the origin is most consistent with serial excitation from reference to target cell, schematized by an excitatory synapse between unit 149-2c and 149-2b.

Lagged troughs were observed as infrequently (4/84) as lagged peaks. Figure 5 illustrates a pair of cells that fired reciprocally and exhibited a lagged correlogram trough. The response averages document the reciprocal firing patterns of these cells during the ramp-and-hold torques. The cross-corre-
The logram is composed of two components: a lagged trough beginning at +6 ms and a small central peak. The lagged trough could not be explained by the refractory period in the target unit’s autocorrelogram and provides evidence for serial synaptic inhibition. The central peak is small but statistically significant. The simplest synaptic circuit explaining these correlogram features is a combination of common synaptic input and serial inhibition. The reference unit (40-5a) was an NPT cell situated 300 μm superficial to unit 39-1a and 10 μm horizontally (tangent to layer V, as judged by radial fiber orientation). Three additional cell pairs demonstrated pure lagged troughs in their correlograms but were poorly modulated with the task.

The presence of common synaptic input to, or serial excitation between, precentral cells with similar firing characteristics makes functional sense, as does a lagged correlogram trough between precentral cells with reciprocal responses. However, several counterintuitive relationships were also observed, as illustrated by the examples in Figs. 6 and 7. Figure 6 shows a pair of cells recorded midway down the bank of the precentral gyrus; their correlogram had a lagged central peak and yet the response averages reveal reciprocal firing patterns. The presence of a lagged central peak (seen in 21/84 significant correlograms) was interpreted as evidence for both common synaptic input and serial excitation (Dickson and Gerstein 1974), shown as a recurrent collateral from reference PT cell 139-1a. This interpretation is supported by further evidence analyzing the time course of these components during movement (Smith 1989). Although the response similarity is negative, the discharge patterns of these cells are not entirely reciprocal. The output target muscles of unit 139-1b were documented by spike-triggered averaging of EMG (Smith and Fetz 2009), which revealed postspike facilitation (PSF) in flexor muscles; unit 139-1a was a PT cell but produced no PSF in any recorded muscles.

Figure 7 illustrates the last correlogram feature type, a central trough (seen in 11/84 significant correlograms), and another example of a “paradoxical” response relationship. Moore et al. (1970) interpreted such correlogram troughs as evidence for internuncial cells that inhibit one cell of the pair while receiving input in common with the other cell, as illustrated. Under these circumstances the target cell will have a lower probability of firing when the reference cell is activated by the common afferents and, conversely, the reference unit will fire less when the target unit is disinhibited. One cell pair had an exceptionally wide central trough (102 ms) that was considered secondary to response coordination (i.e., the trough could be explained by rapid, reciprocal changes in activity evident in the response averages) and this pair was excluded from further analysis. For the pair in Fig. 7 the correlogram trough lasted 34 ms, much briefer than any variation in the response averages. The units fired reciprocally in the flexion direction, but covaried during extension. Compiling correlograms selectively during the flexion and extension phases of torque resulted in too few triggers for meaningful comparison.
The target unit was a PT cell that did not produce PSF in any of the recorded forearm muscles; unit 102-2b is an NPT cell.

Table 1 summarizes the frequency of correlogram feature types found within this sample of 215 cell pairs. Of 84 significant correlograms, 65 (77%) had peaks straddling the origin, consistent with common synaptic input. Of these, 21 had superimposed lagged peaks. Only 8 (10%) had predominantly serial features.

Response similarity and correlogram features

Although 39% of cell pairs had significant cross-correlogram features, 61% of pairs did not. It seems plausible that cells with similar or reciprocal firing properties could be coordinated by synaptic connections, whereas cells firing in an unrelated fashion would be less likely to have synaptic interactions. This hypothesis predicts that pairs with strong covariation in firing properties would have significant correlogram features and cells that fire independently would have flat correlograms.

To test this hypothesis we measured the similarity in response properties of both neurons by the “response similarity,” defined as the correlation coefficient between the smoothed firing rates of the two cells during active wrist torques (see METHODS, Fig. 2): it ranges from -1 to +1 and represents how similarly (positive) or dissimilarly (negative) the two cells fire during the isometric tasks (see METHODS). Numbers near zero represent cells that fire in an unrelated manner. Examples of response similarity values are given in Figs. 4–7 as the “r” numbers between the respective response averages.

The distribution of response similarity values for cell pairs with significant and nonsignificant correlograms is shown in Fig. 8A. The distributions are nearly symmetric around zero, with a mean for all pairs of 0.035 and median of 0.039. Although correlated pairs were out-numbered by uncorrelated pairs, the two distributions are not significantly different. The occurrence of a significant, positive correlogram feature (central peak, lagged central peak, or lagged peak) was no more frequent between cell pairs with positive response similarity than negative (P > 0.25). Similarly, significant, negative correlogram features (central trough or lagged trough) were no more frequent among cell pairs with negative response similarity than positive (P > 0.43). Thus the observation that two cortical cells respond similarly during an active task does not imply that evidence for consistent synaptic interaction will be found in the correlogram.

However, within the population of correlated cell pairs a relationship between the correlogram feature magnitude—which can be taken to represent the strength of synaptic interaction—and response similarity was observed. Figure 8B plots A<sub>N</sub> (the area of the correlogram feature normalized to the algebraic mean of the number of target and reference cell spikes) against response similarity for all significant
correlograms. The graph was divided into four quadrants (I–IV). Cell pairs appearing in quadrant I had positive correlogram peaks and similar response properties, whereas quadrant III represents correlograms with troughs from cells with reciprocal response properties. For these quadrants the correlogram features are functionally consistent with response similarity. In contrast, the cells in quadrants II and IV represent a counterintuitive situation. The regression line is significant ($r = -0.25$, $P = 0.001$, $n = 84$) and the positive slope indicates that stronger response similarity is correlated with larger correlogram features. Quadrant I contains 41 points (49%) and quadrant III contains 9 (11%), representing 60% of significantly correlated cell pairs. The majority of the remaining pairs fell in quadrant II ($n = 28$, 33%); only 6 (7%) occurred in quadrant IV. Each symbol is coded for the six possible combinations of cell types in the pair (see legend); there was no systematic dependence of synaptic interaction or response similarity on cell type.

When analyzed separately by correlogram feature type, no clear relationship between feature type and response similarity was found. Lagged peaks ($n = 4$) were evenly distributed between quadrants I and II (e.g., Fig. 4); i.e., only half of serial, excitatory connections were between cell pairs with similar firing characteristics. The lagged troughs ($n = 4$) all fell within quadrant III; however, in three of the four cell pairs, one cell in each was weakly modulated with the motor task. Central troughs ($n = 11$) were evenly distributed between quadrants III ($n = 5$) and IV ($n = 6$). Finally, although more central peaks and lagged central peaks fell within quadrant I ($n = 39$) than within quadrant II ($n = 26$), the difference was not statistically significant, alone or in combination ($P > 0.10$).

**Duration and strength of synaptic interactions**

The duration and strength of the synaptic interactions were quantified as the width and normalized area of the correlogram feature. The average measures are given in Fig. 9 for each type of feature. The bars further distinguish these values for each of the three possible pairings of PT and NPT cell types (CM was combined with PT). Contrary to the expectation that the morphological differences between pyramidal and nonpyramidal cells might lead to different interaction measures, the parameters were not significantly different for pair types (within each feature type). However, the overall averages (rightmost bars of each category) show significant differences across features. The widths of central peaks and lagged troughs were similar, whereas central troughs were significantly wider and lagged peaks were significantly narrower. The area of central peaks was not significantly different from that for lagged peaks, although central troughs had a larger negative area than that of lagged troughs.

Our cross-correlograms were constructed for all activity, throughout the behavioral task. To document variation in

**FIG. 6.** A PT and CM cell recorded on the same electrode with correlogram exhibiting a lagged central peak and response averages showing largely reciprocal activity. The cells were recorded in the anterior bank of the precentral gyrus. The lagged component of the central peak was interpreted as evidence for serial excitation from unit 139-1a to unit 139-1b, represented as a recurrent collateral from the PT cell.
syntatic interactions during the ramp-and-hold task, we also
compiled cross-correlograms during separate phases of the
task, as well as “joint poststimulus time histograms” (JPSTHs)
(Aertsen et al. 1989) aligned at response onset. Of 73 cell pairs
analyzed with JPSTH, 39 showed no obvious modulation in the
JPSTH during the ramp-and-hold torque trajectories. A third of
the pairs \( n = 25 \) showed a decline in the JPSTH during times
of phasic increases of firing of one or both neurons. This
counterintuitive drop in interactions during an increase in firing
rate was confirmed by selective cross-correlograms. As de-
scribed elsewhere (Smith 1989), changes in components of the
JPSTH and correlograms suggest that serial and common input
mechanisms may be modulated independently.

**TABLE 1. Number of significant correlograms by pair and feature type**

<table>
<thead>
<tr>
<th>Feature</th>
<th>CM–CM</th>
<th>CM–PT</th>
<th>CM–N</th>
<th>PT–PT</th>
<th>PT–N</th>
<th>N–N</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of pairs</td>
<td>12</td>
<td>22</td>
<td>40</td>
<td>15</td>
<td>75</td>
<td>51</td>
<td>215</td>
</tr>
<tr>
<td>Significant correlograms, %</td>
<td>6 (50)</td>
<td>8 (36)</td>
<td>12 (30)</td>
<td>9 (60)</td>
<td>28 (37)</td>
<td>21 (41)</td>
<td>84 (39)</td>
</tr>
<tr>
<td>Central peaks</td>
<td>6</td>
<td>7</td>
<td>7</td>
<td>6</td>
<td>12</td>
<td>12</td>
<td>44</td>
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<tr>
<td>Lagged central peaks</td>
<td>3</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>9</td>
<td>4</td>
<td>21</td>
</tr>
<tr>
<td>Lagged troughs</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Central troughs</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>6</td>
<td>0</td>
<td>11</td>
</tr>
</tbody>
</table>

Entries are number of cell pairs; values in parentheses indicate the percentage of cell pairs that were correlated for each column.

*Strength of synaptic interaction and cell separation*

If the cortex were isotropic with respect to synaptic connec-
tions, the prevalence of synaptic interconnection and the
strength of synchrony between any two cells would decline
inversely with distance to the third power. However, anatom-
cal evidence (Jones and Wise 1977) suggests that corticocor-
tical and thalamocortical connections may be distributed to
motor cortex in mediolateral strips. The width of these strips is
about 0.5 mm and their spatial periodicity is about 0.5–1.0 mm.
Within our study, all cell pairs were recorded within a plane
perpendicular to the central sulcus and thus perpendicular to
the periodic bands of afferent fiber terminations. To determine
This suggests that synaptic interactions, most of which are common synaptic input, are concentrated within 400-μm columns in cortex.

The strength of correlation, measured by $A_N$, was not related to horizontal separation (Fig. 10B). Although the largest correlogram features were found for pairs recorded on a single electrode, $A_N$ assumed a constant value for larger distances. Normalized area remains essentially constant if the two points at 100 μm are excluded. The open circles in this figure represent lagged peaks (positive areas) and lagged troughs (negative areas). The separation of serial inhibitory pairs ranged from 250 to 490 μm; no serial inhibition was observed between pairs recorded on the same electrode.

Laminar location of recorded cells

The synaptic interactions between precentral cells depended in part on the cortical lamina of each cell. To estimate the laminar location of a recorded cell, electrode tracks were superimposed over the appropriate histologic section. Cells were classified as belonging to layers II–III, V, or VI. Of 215 cells, 17 were in layers II–III, 180 were in layer V, and 18 were in layer VI. The preponderance of layer V cells reflects a methodological bias to record as many PT and CM neurons as possible. Larger neurons tended to be sampled preferentially due to the electrodes’ recording bias and the need for long-term stable isolation.

Figure 10A plots a histogram of correlated and uncorrelated pairs as a function of horizontal separation. The pair of bars at 0 separation represents pairs recorded on the same electrode, 51% of which were correlated. The combined prevalence of correlation was higher in the range from 100 to 400 μm than that between pairs on a single electrode. Beyond 400-μm separation this prevalence declines to <25% on average. Separating the data into two ranges at 400 μm, 51% of pairs less than this distance and 25% greater than this were correlated.

whether correlation occurred at a preferred cell separation, the separation for each cell pair was derived from histological reconstructions. We projected each cell to layer V along the orientation of radial fibers and calculated the line integral of distance between the cells along layer V, termed horizontal separation (see METHODS).

Figure 10B plots a histogram of correlated and uncorrelated pairs as a function of horizontal separation. The pair of bars at 0 separation represents pairs recorded on the same electrode, 51% of which were correlated. The combined prevalence of correlation was higher in the range from 100 to 400 μm than that between pairs on a single electrode. Beyond 400-μm separation this prevalence declines to <25% on average. Separating the data into two ranges at 400 μm, 51% of pairs less than this distance and 25% greater than this were correlated.
Table 2 shows the number of correlated pairs listed by the locations of reference and target cell and by the type of correlogram feature. Central peaks were found between 42% of correlated cells within layer V; 91% of these were separated by <600 µm horizontally. Only one of the layer II–III cells shared common inputs with a layer V cell. All correlograms with central troughs were found for neurons within layer V. No significant differences existed between the mean normalized areas of correlogram features for pairs of cells located in the same or different laminae.

**DISCUSSION**

These results elucidate the degree to which synaptic interactions between motor cortex neurons are correlated with their modulation during isometric hand torques. The results augment the body of data on neuronal responses in different regions during this ramp-and-hold torque task (summarized in Fetz et al. 2002). The major types of interactions provide interesting comparisons with results from other studies of motor cortex and from other cortical regions.

**Common synaptic input**

Most of the cell pairs with significant correlogram features (90%) had features that straddled the origin; three fourths had central peaks, consistent with common synaptic input (Moore et al. 1970) or correlated input (Brody 1999a,b). Central peaks are the most prevalent feature found for cell pairs in various cortical regions, including visual cortex (Michalski et al. 1983; Toyama et al. 1981a; T’so et al. 1986) auditory cortex (Dickson and Gerstein 1974; Eggermont 1994; Frostig et al. 1983; Nelson et al. 1992), association area (Noda and Adey 1973), and motor cortex (Allum et al. 1982; Jackson et al. 2003; Kwan et al. 1987). Our results also agree well with a study of synaptic interactions between pairs of precentral cortex neurons determined by STAs of intracellularly recorded membrane potentials (Matsumura et al. 1996). In that study 82% of STAs with features had average synchronous excitatory potentials (ASEPs), which began before the trigger spike. These ASEPs produce central correlogram peaks whose duration typically coincides with the rising edge of the ASEP (Matsumura et al. 1996). The rise time of the ASEPs (18 ± 7 ms) agrees well with the mean width of our central peaks (22 ms).

Two types of mechanisms could produce central peaks: last-order “common input” neurons that send divergent synaptic connections to both of the correlated cells or synchronized input from separate but correlated sources. Direct common input most likely explains the narrower (≈20 ms) central peaks (Nelson et al. 1992). In this case, the number of synchronous
inputs from such common cells can be estimated by comparing the size (i.e., area) of the central peak to that of lagged peaks produced by serial connections. The probability that an afferent fiber that bifurcates to cortical neurons will fire both cells together is the product of the probabilities that an afferent impulse will produce a spike in each cell. If the mean synaptic efficacy of a serial connection is \( \alpha \), the expected contribution of a common input cell to a central correlogram peak of area \( A_N \) would be \( \alpha^2 \). The average value of \( \alpha \) for the serial, excitatory correlograms in this study was 0.032. A common afferent fiber with this efficacy would be expected to produce a central peak with \( A_N = 0.0010 \). The average \( A_N \) for all central peaks was 0.051, roughly 50-fold higher than that expected from a single fiber. This would suggest that about 50 synchronous inputs from common afferents would be expected to mediate the average central peak.

The central peaks could also be produced in part by separate but synchronized inputs (Brody 1999a,b; Lampl et al. 1999). These would be tightly timed, aperiodic synchronous inputs. In monkeys making free reaching movements oscillatory activity in sensorimotor cortex can transiently synchronize large populations of neurons (Baker et al. 1999, 2001; Donoghue et al. 1998; Murthy and Fetz 1992, 1996). Such oscillatory activity does not seem a likely explanation of our central correlogram peaks because careful analysis of the cross- and autocorrelograms of the cells in this study revealed no evidence of periodicity. Moreover, the cortical oscillations entrained cells over wide regions, whereas the central peaks were seen for cells separated by only several millimeters. This suggests that the synchrony described here is not due to periodic oscillations, but rather involves relatively direct common input neurons or aperiodic synchronized inputs.

**Serial excitation**

In this study 5% of the correlograms with significant features showed lagged peaks, indicative of serial excitation. This compares with 8% of STAs that showed pure excitatory postsynaptic potentials (EPSPs) in vivo (Matsumura et al. 1996). The mean width of our lagged correlogram peaks (10 ms) is slightly longer than the mean rise time of the serial EPSPs (6 ± 3 ms). This is consistent with the fact that in cortical neurons, transient depolarizing potentials trigger action potentials not only on their rising edge by direct threshold crossings, but also later, due to delayed crossings produced by a regenerative inward current (cf. Fig. 2 in Fetz et al. 1991; Reyes and Fetz 1993). Later spikes can also be initiated by the superposition of the decaying EPSP and synaptic noise (Fetz and Gustafsson 1983).

The peak area (\( \alpha \)) of our four lagged peaks, normalized to the number of triggers, represented 0.041 above-baseline counts per trigger. Using the transform between EPSPs and the correlogram peaks they produced in vivo (Matsumura et al. 1996), this value would predict that the underlying EPSPs had mean amplitudes of 133 \( \mu \)V. This prediction is smaller than the mean EPSP amplitudes observed in STAs in vivo (226 ± 130 \( \mu \)V) and in vitro (Reyes and Sakmann 1999; Reyes et al. 1998; Thomson et al. 1993a,b), but compatible with the range of values.

Many correlograms (\( n = 21 \)) in this sample showed lagged central peaks, which may represent a combination of recurrent excitation superimposed on common synaptic input (Dickson and Gerstein 1974). Lagged central peaks between PT and NPT cells (\( n = 11 \)) nearly always showed lagged peaks in the direction of PT to NPT cell (\( n = 10 \)). Asymmetric peaks were also observed between PT pairs, including CM cells, but were not seen between NPT pairs. There is anatomical (Ghosh and Porter 1988a) and physiological (Kang et al. 1988; Stefanis and Jasper 1964) evidence for recurrent collaterals from PT cells onto other PT cells in motor cortex.

**Serial inhibition**

Of the correlograms with significant features, 5% showed lagged troughs, indicative of serial inhibition. This compares with 4% of STAs that showed pure serial inhibitory postsynaptic potentials (IPSPs) in vivo (Matsumura et al. 1996). The duration of the four lagged troughs observed in this study averaged 26 ms, which is comparable to the mean duration of the serial IPSPs (25 ± 4 ms). Lagged correlogram troughs with long durations were also found in cat visual cortex neurons (>80 ms) by Toyama et al. (1981b) and in pericruciate cortex (36–60 ms) by Renaud and Kelly (1974). Direct comparison of PSPs and correlograms in motoneurons showed that the duration of the correlogram trough exceeded the rise time of the IPSP but was shorter than the duration of the IPSP (Fetz and Gustafsson 1983).

The relative paucity of evidence for serial inhibition from correlograms seems discrepant with electrophysiological and anatomical studies indicating a predominance of intracortical inhibition within motor cortex. Intracortical inhibition is presumably responsible in part for sculpting the response properties of motor cortex cells. Its importance was demonstrated by the fact that iontophoresis of bicuculline in motor cortex converted responses of precentral neurons from unidirectional responses during limb movements to bidirectional responses (Matsumura et al. 1991). Of the eight morphological types of intracortical neurons (Jones 1975b) only one type—the spiny stellate cell of layer IV—is believed to be excitatory. Thus inhibition should be the predominant serial connection to be found. The lack of correlograms with lagged troughs may be explained by several factors. First is the likely sampling bias against recording such small internuncial cells. Most presumed inhibitory neurons have a relatively symmetric dendritic arborization (closed-field neurons) and a small soma that would make them difficult to record and would bias extracellular recordings to large, open-field pyramidal cells. Second, cross-correlation techniques are less sensitive for detecting inhibition, as suggested by simulation studies (Aertsen and Gerstein 1985). Third, inhibitory troughs may be masked by central peaks produced by concomitant common inputs.

**Synaptic interactions between cells in cortical layers**

The connections revealed by cross-correlation features documented in this and the companion study (Smith and Fetz 2009) are summarized schematically in Fig. 11. The three major types of precentral cells are shown at their laminar locations. Each of the illustrated connections between cells corresponds to an observed correlation. Extracortical inputs, although not specifically identified, are shown to arise from specific thalamic afferent fibers (from nucleus ventralis lateralis) and corticocortical fibers arising from cortex outside area 4.
The cortical cells are grouped into three “columns” of about 500-μm width, in accordance with the range over which neurons were found to receive common inputs. Common inputs were observed to synchronize cells within and across cortical layers, including cells in layers II–III, V, and VI (left column). Pairs of layer V neurons received common inputs (middle column) or were connected in a reciprocal fashion (right column). A similar broad distribution of common inputs has been observed in visual cortex (Douglas and Martin 2004; Toyama et al. 1981a).

Reciprocal collaterals from layer V cells were found to project to other layer V cells and to layer VI cells (middle column). No intracortical excitation was observed to originate from cells outside layer V, suggesting that a major source of intracortical excitation derives from recurrent collaterals of pyramidal cells. In contrast, in visual cortex excitatory connections have been observed from layer III–IV border cells to supragranular cells and from supragranular cells to layer V cells (Komatsu et al. 1988; Thomson and Lamy 2007; Toyama et al. 1981a). Such differences may reflect differences in cortical organization between primary sensory and motor cortices or sampling bias.

Serial inhibitory connections were found between layer VI and layer II–III cells (left column) and between layer II–III and layer V cells (middle column). In visual cortex, layer III–IV border cells have been shown to provide inhibitory connections to supragranular neurons (Komatsu et al. 1988; Thomson and Lamy 2007; Toyama et al. 1981a). Again, such differences may be accounted for by the different methodologies used (intracellular vs. extracellular) or by different synaptic organization of these two cortical regions.

Relation between synaptic interaction and firing properties of precentral cells

This study was designed to address the degree to which the firing properties of motor cortex cells are correlated with their synaptic interactions. As shown in Fig. 8A, the presence or absence of a correlational linkage did not depend on how similarly two cortical units fired during the operant task. In other words one cannot infer functional interactions between two cortical neurons by comparing their response patterns. Obvious as this point may seem, there is a natural and historical tendency to infer such interactions. For example, the correlations between the smoothed firing probability of cells and muscles have been used to infer functional interactions (Houk et al. 1987), although such measures of covariation are task dependent and do not establish causal interactions. At the cortical level we found that even neighboring cells with highly similar response patterns are not necessarily coupled by monosynaptic or common input connections. This suggests a sparse connectivity, in which cells are driven in parallel by separate inputs with similar response patterns. Likewise, two cortical cells with reciprocal response patterns are not necessarily coupled by inhibitory linkages. A relevant caveat is that the prevalence of reciprocal inhibition may have been underestimated because significant correlograms are more difficult to compile for reciprocally activated pairs and for relatively inactive cells.

In contrast, when pairs of precentral cells did have a correlogram feature, its size was significantly related with the similarity in their firing patterns during volitional wrist torques (Fig. 8B). Nevertheless, the relationship is weak because the
effect of unitary EPSPs is a brief statistical increase in the firing probability of the postsynaptic cell (Cope et al. 1987; Moore et al. 1970; Reyes and Fetz 1993). Previous studies have noted qualitative agreement between measures of correlation and response patterns (Allum et al. 1982; Fetz et al. 1991; Georgopoulos et al. 1993). A potential confound in this relationship is the possibility that covariation can influence correlation measures. For example, a measure based on the first recurrence times of spikes in one cell relative to spikes of the second was correlated with the similarity in directional tuning (Georgopoulos et al. 1993), but was also influenced by covariation (Fetz and Shupe 1994; Georgopoulos et al. 1994).

In the present study, the sign and magnitude of correlation features ($A_N$) were significantly related to response similarity (Fig. 8B). Pairs with positive response similarity tended to have larger positive correlogram peaks than those of cells with dissimilar firing patterns. Likewise, pairs with negative similarity had a larger proportion of correlogram troughs than that of pairs with positive similarity. Thus it appears that synaptic interactions do help to shape the response pattern of precentral neurons. However, the magnitudes of individual correlational linkages are relative small, indicating that many neurons must be involved in determining activity during a motor task.

The linear regression in Fig. 8B does not go through the origin, but is shifted upward. This represents the contribution of positive correlation features, which in most cases were central peaks. These correlation peaks appear across the range of response similarity and may be viewed as subserving another functional role: a form of sensorimotor binding or coordination of cells involved in the same behavior. Thus the task-related modulations are separable and appear superimposed on a broader cofacilitation of cells involved in the task.

Conclusions

The present results indicate that the synaptic interactions between motor cortex neurons during voluntary wrist torques are largely consistent with their firing properties. However, the relationship is weak and the scatter is large. This indicates a considerable degree of independence between connections and coactivations. To determine whether this is a consequence of including diverse cell types, the companion paper reexamines the issue for corticomotoneuronal cells, which have correlational linkages with identified target muscles (Smith and Fetz 2009).

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References


SYNAPTIC INTERACTIONS BETWEEN MOTOR CORTEX CELLS


