Direct control of paralysed muscles by cortical neurons

Chet T. Moritz1, Steve I. Perlmutter1 & Eberhard E. Fetz1

A potential treatment for paralysis resulting from spinal cord injury is to route control signals from the brain around the injury by artificial connections. Such signals could then control electrical stimulation of muscles, thereby restoring volitional movement to paralysed limbs1–3. In previously separate experiments, activity of motor cortex neurons related to actual or imagined movements has been used to control computer cursors and robotic arms4–10, and paralysed muscles have been activated by functional electrical stimulation11–13. Here we show that Macaca nemestrina monkeys can directly control stimulation of muscles using the activity of neurons in the motor cortex, thereby restoring goal-directed movements to a transiently paralysed arm. Moreover, neurons could control functional stimulation equally well regardless of any previous association to movement, a finding that considerably expands the source of control signals for brain-machine interfaces. Monkeys learned to use these artificial connections from cortical cells to muscles to generate bidirectional wrist torques, and controlled multiple neuron–muscle pairs simultaneously. Such direct transforms from cortical activity to muscle stimulation could be implemented by autonomous electronic circuitry, creating a relatively natural neuroprosthesis. These results are the first demonstration that direct artificial connections between cortical cells and muscles can compensate for interrupted physiological pathways and restore volitional control of movement to paralysed limbs.

Spinal cord injury impairs neural pathways between the brain and limbs, but spares both the motor cortex and muscles. Recent studies have shown that quadriplegic patients could volitionally modulate activity of neurons in the hand area of the motor cortex, even after several years of paralysis2, and that monkeys could use cortical activity to control a robotic arm to acquire targets5 and feed themselves3. These and other brain–machine interface studies used sophisticated algorithms to decode task-related activity of neural populations and to calculate requisite control parameters for external devices4–10–13. An alternative strategy to restore limb function is to directly connect cortical cell activity to control the stimulation of a patient’s paralysed muscles (Fig. 1a). Here we show that monkeys can learn to use direct artificial connections from arbitrary motor cortex cells to grade stimulation delivered to multiple muscles and restore goal-directed movement to a paralysed arm.

In previous biofeedback studies monkeys rapidly learned to control the discharge rates of newly isolated neurons in the motor cortex to obtain rewards4–13. We used similar operant conditioning techniques for single neurons in the hand and wrist area of the motor cortex of two monkeys (see Methods and Supplementary Information). We tested volitional control of cell activity by displaying smoothed discharge rate as cursor position on a monitor and rewarding the monkeys for maintaining activity within randomly presented high- or low-rate targets. The directional tuning of most cells was also characterized in an isometric two-dimensional wrist target-tracking task. Our experiment, however, used all sufficiently well-isolated cells encountered, with no selection bias for possible association to movement or directional tuning.

Monkeys demonstrated volitional control of the discharge rates of nearly all cells tested within the first 10-min practice session. Although cell activity controlled the cursor directly, monkeys often continued to produce wrist torques during these initial sessions (Supplementary Fig. 1). We then blocked peripheral nerves innervating the wrist muscles with a local anaesthetic (see Methods). Despite loss of motor function and sensory feedback from the innervated forearm, monkeys continued to control the cursor with cell

Figure 1 | Brain-controlled functional electrical stimulation (FES) of muscle. a, Schematic shows cortical cell activity converted to FES during peripheral nerve block. b, Example of motor cortex cell activity controlling FES of paralysed wrist extensors. Extensor (red shading) and centre (grey shading) wrist torque targets were randomly presented. Monkeys learned to modulate smooth cell rate to control proportional muscle stimulation. FES was delivered to muscles EDC and ED4,5 at 50 s−1, with current proportional to cell rate above a stimulation threshold (0.4 mA pps−1 × [cell rate − 16 pps]; ≤10 mA). Here pps indicates pulses per second. c, Histograms of cell rates while acquiring the extensor and centre targets, illustrating cell activity used to successfully control FES. Shading indicates target hold period and horizontal line denotes baseline cell rate.

1Department of Physiology & Biophysics and Washington National Primate Research Center, University of Washington, Seattle, Washington 98195, USA.
activity for 45 out of 46 cells after the nerve block. Supplementary Fig. 1 shows the loss of flexor and extensor torques after injections of local anaesthetic, while the monkey continued to volitionally control the cell activity. The nerve block was confirmed by the inability of the monkeys to perform the two-dimensional torque-tracking task.

We then converted cell activity into proportional stimuli delivered to paralysed muscles, generating functional electrical stimulation (FES). The cursor was now controlled by wrist torque, and the monkeys were rewarded for maintaining FES-evoked torque within peripheral and centre (that is, zero-torque) targets for 0.5–1.0 s. To allow the monkeys to grade contraction force, stimulation current was made linearly proportional to cell rate when the cell discharged above a threshold.

The example in Fig. 1b, c shows a monkey modulating cell activity to generate appropriate torques by controlling FES of paralysed wrist extensor muscles. The monkey learned to increase cell activity to activate the stimulator and acquire the extensor targets, and to maintain activity below the stimulation threshold to relax the muscle and acquire the centre targets. Both monkeys were able to control muscle FES during nerve block and acquire torque targets with 44 out of the 45 cells tested (5 cells from monkey I and 39 from monkey L).

For each cell the monkeys' control improved with practice, as evidenced by more rapid acquisition of targets and fewer errors. Monkeys began using cell activity to control the stimulator almost immediately, and improved substantially during the relatively brief practice sessions with each cell (mean duration 66 min). To quantify this improvement we compared performance during the initial two minutes of practice and during the two-minute period with the highest performance, typically just before task difficulty was increased to probe the limits of FES control. The rate of target acquisition with FES control was over 60% during the two-minute period with the highest performance, while the rate of target acquisition with FES control was just before task difficulty was increased to probe the limits of FES control. The rate of target acquisition with FES control was over 60% during the two-minute period with the highest performance, while the rate of target acquisition with FES control was just before task difficulty was increased to probe the limits of FES control. The rate of target acquisition with FES control was over 60% during the two-minute period with the highest performance, while the rate of target acquisition with FES control was just before task difficulty was increased to probe the limits of FES control.

With continued practice monkeys also learned to control the torque more precisely with cell activity, making fewer target acquisition errors and often acquiring targets on the first attempt. A target acquisition error was defined as triggering the stimulator to acquire the peripheral target when the centre target was displayed. Monkeys made target errors on only 0.8 ± 5.1% of targets during peak performance for each cell compared to 20.7 ± 28.9% of targets at the beginning of practice (P < 0.001; Supplementary Fig. 3). They also made 81% fewer failed attempts to acquire the target during peak performance (0.10 ± 0.31 failed attempts per target) compared to the beginning of practice (0.52 ± 0.93; P < 0.001).

To test whether FES could also be controlled by decreases in cell activity, we set stimulation current to be inversely proportional to cell rate below a threshold for 11 cells. Monkey I learned to control stimulation with this inverse relation just as well as with a positive relation between cell rate and stimulus current (38 cells, some tested in both groups; P > 0.46), acquiring 13.4 ± 3.9 targets per minute and making no errors during peak performance.

The activity of a single cell could also be used to control stimulation of antagonist muscle groups and restore bidirectional movements. Figure 2 shows an example of one cell that controlled stimulation of flexor muscles with high discharge rates and extensor muscles with low rates. The monkey learned to control cell activity and grade contraction force to readily satisfy targets at five different torque levels. The nerve blocks remained very effective, as evidenced by negligible torques produced in either direction when the stimulators were turned off during target presentation (Fig. 2b). Seven cells tested with such bidirectional control performed similarly to cells that controlled only one muscle group, although target acquisition rates were marginally slower (9.8 ± 3.7 targets per min; P = 0.06).

Figure 2 | Brain-controlled FES of multiple muscles restores graded torque in two directions. a, The monkey acquired targets at five levels of flexion-extension (F-E) torque using the activity of a single cell to grade FES delivered to both flexor (FCU) and extensor (ECU and ED4,5) muscles. Flexor FES was proportional to the rate above a threshold (0.6 mA pps⁻¹ × [cell rate − 24 pps]; ≤10 mA); extensor FES was inversely proportional to the cell rate below a second threshold (0.6 mA pps⁻¹ × [cell rate − 12 pps] − cell rate); ≤10 mA). b, Average torques produced to satisfy the five targets during 12 min of practice. With the stimulator off (shaded periods) the monkey could not produce torques greater than 10% of magnitudes used to acquire the targets (blue and red lines), confirming the efficacy of nerve block. c, Histograms of cell rate used to acquire five target levels (coloured boxes at left). Horizontal lines indicate FES thresholds for flexor (blue) and extensor (red) stimulation.
Two neurons control FES. Monkey L simultaneously modulated activity of two neurons, each controlling proportional stimulation of a different muscle group when above a threshold. Monkey L acquired randomly presented flexor (blue), extensor (red) and centre (grey) targets by using cell 1 to stimulate a flexor muscle (FCU; 0.2 mA pps \( \times \) cell rate \(- 34\) pps) and cell 2 to stimulate extensor muscles (ECU and ED4,5; 0.4 mA pps \( \times \) cell rate \(- 12\) pps).

Figure 3 | Cell directional tuning is unrelated to FES control. a, Responses of an untuned and a strongly tuned cell (solid symbols in b and c). The surrounding peri-event histograms show cell activity while matching (shading) each of eight peripheral torque targets in the flexion-extension (F-E) and radial-ulnar (R-U) plane during the unparalysed tracking task (horizontal lines denote baseline cell rates). The radial plot at the centre summarizes cell activity while matching each peripheral target. b, c, Maximum target acquisition rates during direct brain control of cursor (b) and brain-controlled FES (c) plotted as a function of directional tuning strength for cells recorded during the torque-tracking task (\( n = 38 \)). Performance controlling a cursor directly with cell activity was significantly correlated with cell tuning (b; \( r^2 = 0.33, P < 0.001 \)). Subsequent brain-controlled FES performance was uncorrelated with cell tuning (c; \( r^2 = 0.03, P = 0.33 \)).

Subsequent brain-controlled FES (\( r^2 = 0.03, P = 0.33 \); Fig. 3c). For example, with the untuned cell on the left in Fig. 3a the monkey acquired 18.5 targets per minute. The tuned (\( n = 9 \)) and untuned (\( n = 29 \)) cells showed no differences in any measure of FES control (target acquisition rates, errors or failed attempts; \( P > 0.51 \)).

Extending the strategy of direct neural control to more complex movements will require further control signals. As a first step towards this goal, we tested a monkey’s ability to simultaneously control two cell–muscle pairs. Figure 4 shows monkey L using high discharge rates of one cell to control FES of flexor muscles and high rates of a second cell to control extensor muscles. The monkey learned to independently modulate the activity of five cell pairs in order to control antagonist muscles and rapidly acquire bidirectional torque targets at rates similar to single cells (11.6 \( \pm \) 3.8 targets per min, \( P = 0.32 \)).

These findings have several implications for future approaches to neuroprosthetic control. In contrast to the conventional strategy of deriving control signals from the combined activity of a neural population\(^{8,9,10}\), it may prove efficacious to maintain separate signal pathways from cells to muscles. Using direct channels from single cells to specific muscles may provide the brain with more distinguishable outcomes of the cell activity\(^{16}\) and allow innate motor learning mechanisms to help optimize control of the new connections. The ability of the brain to adapt to new but consistent sensorimotor contingencies has been amply documented\(^{17,18}\), and motor cortex can adapt rapidly to learn new motor skills\(^{19,20}\). Motor circuitry can compensate for drastic changes in connectivity, such as surgically
cross-connected nerves controlling wrist flexor and extensor muscles, or targeted reinnervation for control of prosthetic limbs.

Our finding that monkeys could learn to use virtually any motor cortex cell to control muscle stimulation—regardless of the cell’s original relation to wrist movement (Fig. 3c)—suggests another advantage of directly tapping single cell activity. Strategies based on decoding the activity of neural ensembles to obtain movement parameters or muscle activity depend on finding cells that modulate sufficiently with the output variables during actual or imagined movement. Instead, arbitrary cells available on recording arrays could be brought under volitional control using biofeedback, substantially expanding the source of control signals for brain–machine interfaces. This and previous biofeedback studies have shown that even cells with no discernable relation to muscles can be volitionally modulated after brief practice sessions. Issues concerning the use of individual cells and neural populations for prosthetic control are further discussed in Supplementary Information.

The degree of FES control demonstrated here was limited by the relatively brief training time provided by the transient nerve block. Implanted electronic circuitry will enable adaptive learning over much longer times and under more varied conditions. For example, the autonomous ‘Neurochip’ system can discriminate single cell activity and deliver stimulation through days of free behaviour. In several preliminary FES sessions, we confirmed that this system would allow a monkey to trigger stimulation of a paralysed muscle with cell activity and acquire torque targets (Supplementary Fig. 4). Such autonomous low-power circuits could permit subjects to practice continuously with an artificial connection from brain to muscles or the spinal cord, without requiring complex decoding algorithms or robotic arms. Further development of such direct-control strategies may lead to implantable devices that could help restore volitional movements to individuals living with paralysis.

METHODS SUMMARY

Subjects. Two male Macaca nemestrina monkeys participated in the experiments (4–5-yr-old, weight 4.5–6.5 kg). All procedures were approved by the University of Washington Institutional Animal Care and Use Committee.

Recording and paralysis. The activity of single motor cortex cells was recorded using either acute (Monkey i and l) or chronic (Monkey L) electrodes. Each session began by quantifying the response of cells during an isometric, eight-target wrist torque-tracking task. Volitional control of cell activity was confirmed by operantly rewarding acquisition of targets with a cursor controlled by cell rates. Wrist muscles were then paralysed by injecting anaesthetic (3% chloroprocaine or 2% lidocaine, each with 1:100,000 adrenaline) into catheters cross-connected nerves controlling wrist flexor and extensor muscles, or targeted reinnervation for control of prosthetic limbs.

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METHODS

Cortical recording. Sterile surgeries were performed with isoflurane anaesthesia (1–1.5% in 50:50 O2:N2O). All surgeries were followed by a program of analgesics (buprenorphine 0.15 mg kg−1 and ketoprofen 5 mg kg−1) and antibiotics (cephalexin 25 mg kg−1). Each animal was implanted with a cranial recording chamber over the left hand and wrist area of the motor cortex using stereotaxic coordinates (anterior: 13 mm, lateral: 18 mm) to permit cortical recordings29,30. To obtain longer duration cell recordings, monkey I was re-implanted with a chronic electrode array over the left motor cortex. The array of 12 independently movable microwires is fully described elsewhere7. In brief, 50 μm tungsten wires were threaded through individual polyamide guide-tubes in a 2 × 6 array that was anchored to the skull. This array provided stable recordings from the same isolated cell for the duration of an experimental session, and across multiple days for ten cells24,31,32.

Nerve block implant. Reversible paralysis of the right wrist was achieved with one of two nerve block methods. First, catheters were implanted in the brachial plexus, near cords giving rise to the radial, ulnar and median nerves. Epidural catheters (19 gauge, Arrow International) were inserted into the epineurium surrounding each nerve and anchored in place with cyanoacrylate. Second, nerve cuffs with catheters33 were implanted around the median and ulnar nerves in the upper arm. Catheters terminating in the lumen of each Silastic cuff (4 mm inner diameter, 30 mm long) permitted the nerves to be bathed in anaesthetic. Nerves were identified by electrical stimulation, and catheters were tunneled subcutaneously to exit the skin on the upper back and sealed with an injection port. Thirty-one cells controlled FES during nerve blocks induced by the catheter method, and the remaining 13 cells during blocks induced by cuffs.

Experimental procedures. The monkey sat with his right elbow and hand immobilized by padded splints while a transducer measured the flexion-extension (F-E) and radial-ulnar (R-U) torques produced about the wrist (see Fig. 1a). To receive an appesacuse reward, the monkey maintained wrist torque within a centre target (zero torque) followed by one of eight peripheral targets specifying different combinations of F-E and R-U torque (average magnitude 0.13 ± 0.01 nM). Isolated cell activity was discriminated online using template-matching software (Alpha Omega MSD). Subsequently, cell activity controlled cursor movement in one dimension. Interspike intervals were averaged over a 0.5-s sliding window to create a continuous signal for cursor position (and later FES control). If the cell was directionally tuned, targets were aligned with its preferred direction. For untuned cells or cells without tuning information (that is, cells isolated after nerve block), target errors are reported as the percentage of centre targets presented. A failed attempt to reach a target. A target acquisition error was counted when the monkey activated the stimulator while the centre target was on the screen, resulting in sufficient torque to satisfy any peripheral target had it been presented. Target errors are reported as the percentage of centre targets presented. A failed attempt was counted whenever the monkey briefly acquired a peripheral torque target but did not satisfy the required hold time. A Student’s t-test was used to compare average torques during graded FES control. Otherwise, the non-parametric rank sum test was used for all comparisons as at least one data set for each remaining comparison failed the Lilliefors test for normality. All reported values are means ± s.d.