Contralesional Brain–Computer Interface Control of a Powered Exoskeleton for Motor Recovery in Chronic Stroke Survivors

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Background and Purpose—There are few effective therapies to achieve functional recovery from motor-related disabilities affecting the upper limb after stroke. This feasibility study tested whether a powered exoskeleton driven by a brain–computer interface (BCI), using neural activity from the unaffected cortical hemisphere, could affect motor recovery in chronic hemiparetic stroke survivors. This novel system was designed and configured for a home-based setting to test the feasibility of BCI-driven neurorehabilitation in outpatient environments.

Methods—Ten chronic hemiparetic stroke survivors with moderate-to-severe upper-limb motor impairment (mean Action Research Arm Test=13.4) used a powered exoskeleton that opened and closed the affected hand using spectral power from electroencephalographic signals from the unaffected hemisphere associated with imagined hand movements of the paretic limb. Patients used the system at home for 12 weeks. Motor function was evaluated before, during, and after the treatment.

Results—Across patients, our BCI-driven approach resulted in a statistically significant average increase of 6.2 points in the Action Research Arm Test. This behavioral improvement significantly correlated with improvements in BCI control. Secondary outcomes of grasp strength, Motricity Index, and the Canadian Occupational Performance Measure also significantly improved.

Conclusions—The findings demonstrate the therapeutic potential of a BCI-driven neurorehabilitation approach using the unaffected hemisphere in this uncontrolled sample of chronic stroke survivors. They also demonstrate that BCI-driven neurorehabilitation can be effectively delivered in the home environment, thus increasing the probability of future clinical translation.

Clinical Trial Registration—URL: http://www.clinicaltrials.gov. Unique identifier: NCT02552368.

Key Words: arm • brain–computer interface • hand • rehabilitation • stroke

A significant challenge in the treatment of stroke survivors is the rehabilitation of chronic motor disabilities. Although behavioral therapies such as constraint-induced movement therapy1 or robot-aided sensorimotor stimulation2 can improve upper-limb motor function, they require some level of peripheral motor function to engage with the therapy. This residual function is variable across patients and absent in the setting of complete hemiplegia. An alternative to behavioral therapies is to engage with the patient’s central nervous system directly. Specifically, a brain–computer interface (BCI) system can measure movement-related signals from the central nervous system and provide meaningful feedback to the central nervous system to direct plasticity.

BCIs have recently emerged as novel and potentially powerful tools to restore function in chronic stroke survivors.3 Early results present promising demonstrations that BCI-controlled orthoses or functional electric stimulators can lead to improvements in motor function in chronic stroke survivors.3–5 These stroke-specific BCI systems for rehabilitation have focused on signals stemming from perilesional cortex, contralateral to the...
affected hand for BCI control. Because the ability to modulate perilesional cortical activity decreases with increasing cortical damage, it may be particularly important for neurorehabilitation systems to focus on the ipsilateral, contralesional cortex in those patients who are most severely affected.

Although movement-related neural activity occurs in the ipsilateral and the contralateral cortices, the role of the unaffected hemisphere in stroke recovery is uncertain. Specifically, decreases in contralesional activity are associated with optimal recovery in some studies. Other studies show that increases in contralesional activity may be related to motor recovery. Particular in patients with incomplete recovery. As motor recovery is inversely correlated with the extent of corticospinal tract transection, we hypothesized that using contralesional hemisphere activity to drive a BCI-controlled exoskeleton may lead to functional improvements. Previously, we demonstrated that chronic stroke survivors can control BCIs using electroencephalographic (EEG) signals from the contralesional hemisphere associated with the intention to move the affected limb. However, it was uncertain whether emphasizing the relationship between activation of ipsilateral cortex and resultant sensory feedback would be beneficial.

This feasibility study tested an EEG-BCI system that used signals related to affected hand motor imagery, recorded from the unaffected hemisphere, to control the affected hand via a powered exoskeleton. This study is the first to specifically focus on the unaffected hemisphere with a BCI rehabilitation system and the first to provide BCI-driven therapy in the patients’ homes. This setting is important because it increases the likelihood that this approach can be scaled more widely across the stroke-affected population.

Methods
To determine whether a BCI-controlled exoskeleton using EEG signals from the unaffected hemisphere can lead to functional rehabilitation, we created a novel home-based system called the IpsiHand. We then examined whether a 12-week training period led to functional improvements in chronic, hemiparetic stroke survivors.

Patient Characteristics
Ten chronic hemiparetic stroke survivors with moderate-to-severe upper-limb hemiparesis, enrolled at least 6 months after first-time hemispheric stroke, completed the study. Because motor recovery plateaus after 3 months, the study was designed as a self-controlled study comparing motor function before and after the intervention to establish the feasibility of the BCI-driven therapy studied. The Table contains patient demographics and baseline motor function. The online-only Data Supplement contains detailed inclusion and exclusion criteria. Moderate-to-severely impaired patients were specifically targeted because they are less likely to recover through other methods and therefore require an alternative rehabilitation strategy, such as a BCI. The Washington University School of Medicine Institutional Review Board approved the study protocol, and all patients provided written informed consent.

BCI System Design
The BCI system (Figure 1A) combined a novel powered exoskeleton with a commercial EEG amplifier and active electrodes. The exoskeleton opened and closed the patient’s hand in a 3-finger pinch grip (1 degree of freedom). A detailed description of the system is contained in the Methods in the online-only Data Supplement. Consistent with our previous work, the system used spectral power changes to control hand position. Because stroke patients typically have difficulty extending their extremities, BCI control associated motor imagery with opening the affected hand. Each trial began with the hand fully closed, and spectral power at the control feature was used to update the hand position, providing visual and proprioceptive feedback. During rest trials, patients were instructed to try to keep the exoskeleton closed by imagining that they were resting. During movement trials, patients were instructed to try to open their hand via motor imagery.

EEG Screening
After meeting the inclusion criteria, patients underwent an EEG screening protocol to ensure that a consistent control signal was present for device control. Each patient completed 3 separate screenings to assess the stability of potential BCI control signals. EEG electrodes were applied by a trained biomedical engineer, and EEG signals were collected while patients performed a visually cued motor screening task consisting of trials of (1) rest, (2) unaffected hand movements, (3) affected hand motor imagery, and (4) bilateral motor imagery. Spectral power, or the power in the EEG signal as a function of frequency, was calculated using an autoregressive spectral estimation method. The coefficient of determination ($r^2$), the percent of variance in spectral power that was accounted for by the difference between affected hand motor imagery and rest trials, was calculated for each channel and frequency. After completing 3 EEG screenings, the EEG data were examined for the presence of consistent spectral power changes during affected hand motor imagery. BCI control features were required to be associated with imagined movements of the affected hand and located in unaffected hemisphere motor regions. These sessions were not designed to achieve BCI mastery but to identify patients with consistent cortical activations (ie, $\mu$ [8–12 Hz] or $\beta$ [12–30 Hz] power decreases) in at least 2 of 3 sessions. The feature in the unaffected hemisphere with the strongest $r^2$ value was chosen as the patient-specific BCI control feature. Patients without consistent spectral power changes were unable to continue in the study.

Outcome Measures
The primary outcome measure was the Action Research Arm Test (ARAT). Secondary outcome measures included: (1) the Canadian Occupational Performance Measure, (2) the Motricity Index, (3) the modified Ashworth Scale at the elbow joint, (4) grip strength, (5) pinch strength, and (6) the active range of motion (AROM) at the metacarpophalangeal joint of digits 2 to 5. As this study was the first to use a BCI system for stroke rehabilitation in the home setting, we measured the BCI control quality by comparing the topographies of spectral power changes in the laboratory and home-based sessions. We assessed compliance by recording the total number of days and time that each patient used the system.

Study Protocol
The study timeline is shown in Figure 1B. After completing the EEG screenings, patients completed 2 pretherapy motor evaluations in which all primary and secondary outcome measures were measured by an occupational therapist. On these days, the exoskeleton was also fit to the patient’s hand. In addition, patients and their caregivers were trained to use the system. This included (1) donning the exoskeleton and EEG cap, (2) examining the EEG readouts to verify that physiological signals were collected, (3) software operation, and (4) system maintenance. After the baseline motor evaluations and training, each patient was sent home with a BCI system to complete 12 weeks of therapy. Patients were instructed to use the BCI system on a minimum of 5 days per week. Patients completed 1 to 12 10-minute runs of the BCI task per day depending on their stamina and time constraints. At 2-week intervals, patients came to the laboratory for follow-up motor evaluations consisting of the ARAT and Canadian Occupational Performance Measure. At these follow-up sessions and as needed, an occupational therapist or a biomedical engineer communicated with the patients to ensure compliance with the study.
answer questions about the device, fix any malfunctions, and discuss EEG signal quality, which was assessed regularly by a biomedical engineer. After 12 weeks, patients were again tested on all primary and secondary outcome measures. Different occupational therapists collected baseline and completion outcome measures, and all occupational therapists were blinded to observed EEG changes.

### Table. Patient Characteristics and ARAT Scores

| Patient | Age, y | Time Post-Stroke, mo | Hand Dominance | Clinical Cause/Location | Affected UE | Baseline ARAT | Completion ARAT | ARAT Change |
|---------|--------|----------------------|----------------|-------------------------|-------------|---------------|----------------|-------------|
| 1       | 63     | 49                   | L              | L Ischemic CVA          | R           | 16.5          | 29             | 12.5        |
| 2       | 41     | 18                   | L              | RICA/MCA Dissection leading to a R basal ganglia/ internal capsule stroke | L           | 6.5           | 7              | 0.5         |
| 3       | 72     | 7                    | R              | L Hemorrhagic CVA       | R           | 4             | 12             | 8           |
| 4       | 57     | 29                   | R              | L Thalamic Hemorrhage   | R           | 10            | 16             | 6           |
| 5       | 65     | 12                   | R              | L Periventricular cystic encephalomalacia | R           | 32            | 34             | 2           |
| 6       | 67     | 283                  | R              | R Ischemic CVA          | L           | 15            | 21             | 6           |
| 7       | 62     | 35                   | R              | R Ischemic CVA          | L           | 5             | 6              | 1           |
| 8       | 48     | 6                    | R              | R Ischemic MCA CVA      | L           | 5             | 12             | 7           |
| 9       | 46     | 42                   | R              | L Ischemic CVA          | R           | 29.5          | 43             | 13.5        |
| 10      | 65     | 255                  | R              | R AVM                   | L           | 10.5          | 16             | 5.5         |
| Mean    | 58.6   | 73.6                 |                |                          |             | 13.4          | 19.6           | 6.20        |
| Median  | 62.5   | 32                   |                |                          |             | 10.25         | 16             | 6.00        |
| SD      | 10.3   | 104.2                |                |                          |             | 10.1          | 12.2           | 3.81        |

ARAT indicates Action Research Arm Test; AVM, arteriovenous malformation; CVA, cerebrovascular accident; ICA, internal carotid artery; L, left; MCA, middle cerebral artery; R, right; and UE, upper extremity.

**Figure 1.** Study methodology. **A**, The exoskeleton used attached to a patient’s affected hand via straps on the forearm, palm of the hand, and intermediate phalanges of the index and middle finger, whereas the thumb was held stationary. The exoskeleton was controlled by a microprocessor in the forearm assembly that processed electroencephalographic (EEG) signals. A linear actuator drove hand movements in a 3-finger pinch grip based on the decoded EEG. **B**, The study tested whether training with the brain–computer interface (BCI)–controlled exoskeleton would lead to functional improvements. Patients that met the inclusion criteria completed 3 EEG screenings. Patients with consistent movement-related EEG activations then completed baseline motor evaluations and BCI system training. Finally, patients completed a 12-wk home-based BCI protocol with follow-up motor evaluations at 2-wk intervals.
Results

Ten patients completed the study. Patient characteristics are summarized in the Table, and the online-only Data Supplement contains a detailed description of patient recruitment. In short, of the 22 patients who completed EEG screenings, 18 (81%) were suitable for further BCI therapy, 13 (59%) began the therapy, and 10 (45%) completed the study. The drop off was because of a variety of causes, including unrelated medical diagnoses, inability to comply with the time commitment, and poor orthosis fit.

BCI Control

After initial training, patients and their caregivers were able to apply EEG electrodes in the home setting to record physiological EEG signals. Figure 2 shows exemplary movement-related EEG activity observed in the laboratory and while at home. The patient demonstrated bilateral μ- and β-band power decreases in both settings. Furthermore, the patient had very similar spatial and spectral patterns of movement-related EEG activity during both sessions. The significant decrease in power during motor imagery in the BCI control task led to a high level of accuracy with discriminable patterns of exoskeleton movement during rest and motor imagery.

Because our hypothesis focused on the contralesional hemisphere, the features used to drive the BCI system were from electrodes over the contralesional motor cortex. Movement-related EEG activations were also observed from the ipsilesional hemisphere in 8 of the 10 patients. Although the frequency used for BCI control varied across patients, all BCI control features were μ- and β-band power suppressions, also referred to as event-related desynchronization.22 Patients used the device on 37 to 72 days. Patients performed 74 to 465 10-minute runs of the BCI task for a total of 740 to 4650 minutes of online BCI control in addition to the daily screening task. Details of the patient-specific BCI control are included in the online-only Data Supplement.

Functional Outcomes

The 2 baseline motor assessments were averaged to determine each patient’s baseline motor function. ARAT changes throughout the study protocol are shown in Figure 3A. Patients had a statistically significant mean ARAT increase of 6.2 points. Importantly, 5.7 points has been estimated to represent the minimal clinically important difference in chronic stroke survivors.23 Specifically, 6 of the 10 patients...
had ARAT improvements above this level. In addition to this per-protocol analysis, a significant increase in ARAT score was also found using an intention-to-treat analysis as described in the online-only Data Supplement. Grasp strength, Motricity Index, the grip and grasp ARAT subscores, and Canadian Occupational Performance Measure performance and satisfaction ratings were also significantly increased after therapy, whereas pinch strength, AROM, and the pinch and gross ARAT subscores were not changed. Figure 4 and Table II in the online-only Data Supplement summarize changes across outcomes. Other than minor fatigue, no negative effects were observed.

Neurophysiological Correlates
Across patients, there was a significant correlation between the change in ARAT score and the change in BCI accuracy (defined as the difference between the hand position in the movement and rest trials) per BCI task run (Figure 3B; Spearman $r=0.75$, $P=0.013$). There was not a significant relationship between the change in ARAT score and the total device usage time (Figure 3C; Spearman $r=0.47$, $P=0.17$).

Finally, we sought to determine whether there was a relationship between ARAT and EEG changes (Figure 5). There was a trend toward a positive relationship between ARAT
score changes and the change in the EEG modulation per run of the BCI task at the location and frequency used for BCI control and in a site in the contralateral motor cortex (BCI control feature: Spearman $r=0.48$, $P=0.16$, contralateral motor cortex: Spearman $r=0.62$, $P=0.06$).

**Discussion**

This study provides evidence for the potential role of the unaffected hemisphere in rehabilitation via a BCI-controlled exoskeleton. Specifically, patients had an average ARAT improvement surpassing the minimal clinically important difference. In addition, improvements were observed in some, but not all, objective secondary measures of function. Although pinch strength, AROM, and the ARAT pinch sub-component did not change, these measures are less sensitive in more severely impaired patients and were likely affected by a qualitative increase in spasticity observed, particularly in patients who had received botox 90 to 120 days before study onset. Furthermore, the grasp and grip ARAT subcomponents and grip strength, which all involve distal hand function, significantly improved. It is uncertain whether observed improvements in general distal hand function that did not localize to pinch were because of the poor spatial specificity of EEG or the sensitivity of pinch-specific subcomponents. Finally, we also observed statistically significant increases in a self-scored subjective measure of each patient’s use of their affected arm in functional tasks (Canadian Occupational Performance Measure). These findings build on previous evidence that BCI-controlled rehabilitation systems can facilitate motor recovery. There are several features that distinguish this work from previous studies. First, this study was the first to focus exclusively on using the unaffected hemisphere in a BCI rehabilitation system. Second, the BCI drove the velocity of the exoskeleton, providing a closer temporal pairing between brain activity and proprioceptive feedback than previous systems.

The choice of a BCI control signal for poststroke motor rehabilitation requires careful consideration, particularly given the conflicting evidence on the unaffected hemisphere after stroke. By pairing cortical activations with peripheral feedback, we hypothesized that we would induce plasticity in the remaining (ipsilateral) central nervous system pathways. As noted, there was a significant relationship between the change in ARAT scores and the rate of change in BCI control accuracy that could not be explained by the volume of device use. Further, there was a trend toward a significant relationship between the rate of change in EEG activity and ARAT score specific to the bilateral motor system, but not in the frontal lobe or at task-irrelevant frequencies. Therefore, although what can be asserted from a mechanistic standpoint is somewhat limited, the results indicate that the choice of a BCI control feature in the unaffected hemisphere may have played an important role in the benefits of the intervention.
There are many potential explanations that could account for the functional improvements observed. Specifically, although postrecovery increases in activity have been found in both the affected and unaffected hemispheres,16,24,26,29 the reorganization of interhemispheric connectivity between the contralesional and ipsilesional motor cortices may also play a role in functional recovery.17,28 Further studies designed to better define the mechanism of action will be beneficial to better understand the characteristics of patients who will benefit optimally from BCIs controlled from the unaffected hemisphere. Because the integrity of the ipsilesional corticospinal tract is strongly correlated with motor recovery,17 we would hypothesize that the corticospinal tract integrity is essential in determining what role the contralesional hemisphere will play in recovery. Specifically, in patients with the greatest corticospinal tract damage, we would expect recovery to require an alternative pathway, such as fibers descending ipsilateral to the contralesional motor cortex.

This study was also unique in that the system was used in the home setting without daily oversight. Traditional BCI systems for rehabilitation have been used in a laboratory setting with trained experts operating them.4,4 The ability to provide therapy in a patient’s home without constant supervision would likely reduce the cost of therapy, increase the time of therapy, and give patients flexibility in scheduling therapy. For this approach to achieve large-scale implementation, several practical aspects will need to be addressed, including building the system in a cost-effective fashion, optimizing the orthosis and EEG headset design for enhanced user experience and compliance, and integrating the hardware and software to enable seamless remote maintenance and minimize the need for EEG quality checks.

There are also several limitations to note. Because of the home-based setting, it was impossible to ensure that data were free from artifacts. Although the majority of patients had good-quality EEG recordings in the majority of sessions, a few patients met this standard in <50% of sessions. In addition, because the study sample is small in size and was restricted to those with enough motivation to complete the study protocol, the scope and generalizability of the results is uncertain. Also, pinch strength, all Motricity Index subcomponents, ARAT pinch and gross subcomponents, and AROM did not improve. Whether this was because of the poorer sensitivity of these subcomponents combined with the small sample size, the poor spatial resolution of the EEG signals used, or a limitation of the therapy is uncertain. Finally, the study was uncontrolled. Previous work has shown ARAT improvements can be achieved in chronic stroke patients after interventions such as constraint-induced movement therapy or standard physical therapy,30 but patients in these studies began with a much higher baseline ARAT score than the current cohort. Also of note, while shorter in duration (2 weeks), a randomized controlled trial of a BCI-controlled hand orthosis in patients with a similar baseline motor function showed no improvement in a control group receiving a sham therapy.4 Taken together, there remains an open question of whether more severely affected chronic stroke patients benefit from a BCI intervention exclusively versus prolonged physical therapy; a question that will ultimately be answered with a randomized clinical trial. However, this work provides important early evidence that training with a BCI-driven orthosis can be implemented in the home environment and is associated with a meaningful functional improvement.

Conclusions

This feasibility study shows a statistically significant and clinically meaningful improvement in the motor function of chronic stroke survivors after using a home-based BCI-controlled exoskeleton. The use of control features in the contralesional hemisphere shows evidence of the potential relevance of the unaffected hemisphere for functional rehabilitation. Collectively, although this study represents an important step toward developing and translating BCI-driven rehabilitation protocols for chronic stroke survivors, the effectiveness of BCI-driven therapies must be proven in large randomized controlled trials before full acceptance.

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Disclosures

Dr Bundy, Dr Schalk, R. Coker, Dr Moran, and Dr Leuthardt own stock in Neurolutions, Inc. Study data were reviewed by an unaffiliated neurologist before submission as part of a comprehensive conflict of interest management plan. The other authors report no conflicts.

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SUPPLEMENTAL MATERIAL

Contralesional Brain-Computer Interface Control of a Powered Exoskeleton for Motor Recovery in Chronic Stroke Survivors

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Supplemental Methods

Inclusion/Exclusion Criteria

All patients were chronic, hemiparetic stroke survivors, defined as at least 6 months post first-time stroke. Motor recovery had plateaued and any standard rehabilitation therapy had been discontinued. Specific inclusion criteria consisted of moderate to severe impairment of the upper extremity, limited spasticity (Modified Ashworth score of 1+ or less), full passive range of motion of the affected elbow, wrist, and digits, and normal sensation of the affected upper extremity. As movement of the hand during the trial was completed by the mechanical orthosis, there was no baseline level of active motor control at any joint in the upper extremity required for participation. Full passive range of motion was required, however, in order to ensure that the orthosis could adequately drive hand movements. Exclusion criteria included severe visual impairment, cognitive impairment (8 or more on the Short Blessed Test), botox injections in the affected upper extremity for spasticity management in the prior 3 months, severe aphasia, ataxia, and unilateral neglect.

BCI System Design

The hand component of the exoskeleton was connected to a forearm assembly, which housed a controller board with a microprocessor, motor driver, and touchscreen display for user interface. The exoskeleton was attached to the patient’s hand by straps around the forearm, palm of the hand, and the intermediate phalanges of the index and middle fingers. Because the system was designed for daily use by patients, a limited montage of electrode locations (F3, F4, T7, C3, Cz, C4, T8, Pz) was used. EEG signals were collected using commercially available g.LadyBird active electrodes system and a commercially available g.Mobilab+ EEG amplifier (g.Tec, Graz, Austria). Custom software was written to receive and buffer EEG signals, perform signal processing, and control the position of the exoskeleton. Additionally, the software provided instructions for the patients, received touchscreen inputs to start and stop sessions, and included a display of raw EEG signals to allow patients and researchers to verify that physiologic signals were being captured.

Screening Task

The screening task used to assess spectral power changes associated with motor imagery of the affected hand consisted of 8-second trials of: 1) rest, 2) unaffected hand movement, 3) affected hand motor imagery, and 4) bilateral motor imagery. Each run consisted of 12 trials of each condition and 4 runs were completed in each session for a total of 48 trials of each condition. Data from the screening session was analyzed offline by re-referencing EEG signals to the common average and using an autoregressive method for spectral power estimation known as the Maximum Entropy Method (MEM) to calculate spectral power in 1Hz bins from 1 - 50Hz using 500 msec sliding windows. Following the screening task, a single calibration run (30 affected hand motor imagery trials and 30 rest trials) was performed and served to validate the chosen BCI control feature.

BCI Control Sessions

During online BCI control sessions, EEG signals were re-referenced to the common average and spectral analysis was performed in 1 Hz bins on 500 msec windows of EEG data shifted by 125 msec per window using the MEM algorithm1. After each 500 msec
window was collected, the spectral power at the control feature was used to update the
glove position as described by equation S1:

\[ Y(t) = Y(t-1) + Gain \frac{(X(t) - \mu_{\text{Rest}} - \text{Bias})}{\sigma_{\text{Rest}}} \text{sign}(\mu_{\text{Move}} - \mu_{\text{Rest}}) \]  

(\text{S1})

where \( Y(t) \) is the current glove position constrained to the 0-100% range, \( Y(t-1) \) is the
previous glove position, \( X(t) \) is the current value of the BCI control feature, \( \mu_{\text{Rest}} \) and \( \mu_{\text{Move}} \)
are the means of the BCI control feature during the motor imagery and rest trials, \( \sigma_{\text{Rest}} \) is
the standard deviation of the BCI control feature during the rest trials, \( Gain \) is a gain term
controlling the speed of the movement, and \( \text{Bias} \) is a bias term designed to improve the
ability to discriminate rest periods.

Each run of the BCI control task consisted of 30 rest trials and 30 movement trials.
Each trial was 8 seconds in duration. During rest trials, patients were instructed to try to
keep their hand closed by imagining that they were resting and during movement trials,
they were instructed to try to open the exoskeleton by performing motor imagery. During
control, both visual and proprioceptive feedback of the current hand position was provided
by the exoskeleton. Visually, position was displayed on the touchscreen attached to the
patient’s forearm in the form of a moving bar. Simultaneously, the actuator on the
exoskeleton opened and closed the patient’s hand based upon the spectral power from the
BCI control feature. Patient usage data, including raw EEG signals and the corresponding
hand position, were stored for later analysis.

**Outcome Measures**

The primary outcome measure was the Action Research Arm Test (ARAT). The ARAT
is a 57-point test designed to assess specific changes in upper limb function with sub-
components for grasp, grip, pinch, and gross motor movement\(^2,3\). The ARAT is a standardized
clinical test of arm and hand function used worldwide to quantify post stroke motor deficits in
humans. This test has been validated across numerous studies\(^3\)\(^-\)\(^5\) and found to be equally as
sensitive to other commonly used tools such as the Fugl Meyer Assessment\(^6\). The Canadian
Occupational Performance Measure (COPM) is an evidence-based outcome measure
designed to capture a client’s self-perception of performance in 5 patient-identified tasks
over time\(^7\). At study onset, patients identified 5 functional activities that they wanted to
perform more independently or with greater ease. COPM measurements consisted of a
semi-structured interview in which patients self-rated their performance and satisfaction
with each activity on an ordinal scale from 1 to 10. The Motricity Index provides an overall
indication of a patient’s limb impairment by grading pinch, shoulder abduction, and elbow
flexion on an ordinal scale from 0-5 and reweighting the scale based upon the difficulty
experienced by patients in progressing from one grade to the next\(^8\). Each joint is assigned a
weighted score between 0 and 33 and the scores for the three joints are summed to
produce a score between 0 and 100 for the affected upper limb. The modified Ashworth
Scale measured spasticity on an ordinal scale from 0-4 with an additional intermediate
level (1+) to make the scale more discrete\(^9\). Gross grasp grip strength and three-finger
pinch grip strength were measured using dynamometers. Finally, active range of motion at
the metacarpophalangeal joints of the affected hand was measured relative to full
extension with a goniometer using standard protocols. Positive values for active range of
motion indicated a final position that was in flexion relative to full extension and negative values indicated a final position that was in hyperextension relative to full extension.

**Study Protocol**

Throughout the 12-week study period, patients were instructed to use the BCI system at a minimum level of 5 days per week. On each day, patients began by donning the EEG electrodes. Patients then completed a calibration task in which EEG signals were stored during 90 seconds of rest during which patients were instructed to remain still and 30 trials each of affected hand motor imagery and rest. After completing the calibration task, patients completed one or more runs of a BCI control task with each run consisting of 30 trials in which patients were instructed to attempt to use their EEG activity to open the exoskeleton by performing motor imagery and 30 trials in which patients were instructed to try to keep the exoskeleton in a closed position by resting. Each run of the BCI control task lasted about 10 minutes, patients completed 1-12 runs per day based upon their stamina and other time constraints. The 10-minute duration for each run of the BCI task was chosen as a realistic time period for patients to continuously focus. While patients varied quite a bit in the total time of use per day, because each day required the subjects to don the device and perform the calibration task prior to beginning BCI control and to remove and clean the system after a session, even completion of a single 10 minute run of the BCI control task required at least 40 minutes of total time to complete. Data from each calibration and control run was stored on the system and patients were instructed to maintain a log of their daily usage. Additionally, a wireless hotspot was used to upload anonymized EEG data to an online server. Data was analyzed by an experimenter and used to confirm that physiologic signals were recorded in order to provide feedback to patients about proper system usage.

**Supplemental Data**

**Patient Characteristics**

During the study, 23 patients were enrolled and 22 patients completed all 3 EEG screening sessions. 19 of the 22 patients demonstrated consistent movement-related EEG activity from the unaffected hemisphere ipsilateral to the affected hand. One patient demonstrated activations only contralateral to the affected hand, and two did not demonstrate consistent movement-related EEG activity. Of the 19 potential candidates that successfully completed the EEG screening, six patients did not continue with the study. This was due to the following reasons: 1) impaired cognitive understanding of the system that would have limited the ability to perform the necessary study procedures (1 patient), 2) the exoskeleton did not fit the patient's hand (1 patient), 3) conflicting personal commitments limiting regular usage (2 patients), and 4) health conditions that prohibited consistent use (2 patients). Therefore, 13 patients were eventually sent home with a BCI-driven exoskeleton system. During the study, three patients failed to comply with the study protocol by not utilizing the system at least five days per week and were discontinued from the study. Two of these patients were withdrawn due to an inability to meet the time commitments of the continued device usage and study visits. The third patient was withdrawn because of an unexpected move out of state. Because this study was designed to examine whether training with an powered exoskeleton driven through BCI control from the unaffected hemisphere could lead to functional improvements, data was only analyzed
from the 10 patients who complied with the study for the full 12-week period. While the study specifically focused on using the unaffected hemisphere to drive the BCI system, 8 of the 10 patients also demonstrated consistent movement-related spectral power changes in the ipsilesional hemisphere in addition to the contralesional hemisphere. Where possible the location and type of lesion were collected from patient medical records and are recorded in Table 1 of the main manuscript.

**BCI System Usage**

Supplemental Table I contains information describing the control features used by each patient, the number of runs of the BCI task performed by patients in their homes, and the characteristics of BCI control. Given the home-based context of non-expert electrode application and less controlled noisy environments when the system was being used during this study, careful attention was required when comparing BCI performance or EEG activity to metrics of motor recovery. Specifically, it was important to ensure that experimental runs without physiologic activity were excluded. Therefore, only BCI control runs with significant (p<0.01) \( r^2 \) values indicating differences in EEG activity between movement and rest were included for analysis of the relationship between ARAT changes and BCI performance and EEG activity. While over 50% of the BCI control runs were included in most patients, in a few patients (patients 3, 6, and 9), a larger percentage of BCI control runs were excluded.

**Motor Function Changes**

A detailed description of all outcome scores is shown in Supplemental Table II. At study onset, patients demonstrated moderate to severe motor impairments with ARAT scores ranging from 4-32. Similarly the patients had very low pinch strength scores, Motricity index scores, and while patients could generally perform flexion movements, they struggled to open their hand with no patient able to complete an extension movement to full extension. After the study there were significant (p<0.05) improvements in ARAT score, the grasp and grip subcomponents of ARAT score, Motricity index, grasp strength, and both the performance and satisfaction scores on the COPM.

As described in the manuscript, to establish the potential for BCI training to lead to functional improvements, a per-protocol analysis was used as the primary analysis. While it was not possible to collect completion data for the 3 patients that failed to complete the 12-week study period due to poor compliance, an intention-to-treat analysis was performed using the last ARAT score collected. Across the 13 patients sent home with a device, we observed a mean and median ARAT change of 5 and 5.5 points respectively which was highly significant (p=0.002).
### Supplemental Table I. BCI control features used and characteristics of home-based BCI usage.

| Patient | Affected UE | BCI Control Channel | BCI Control Frequency | Number of BCI Sessions (Days) | Number of BCI Runs | Number of BCI Runs Analyzed | Percent of BCI Runs Analyzed |
|---------|-------------|---------------------|-----------------------|------------------------------|-------------------|-----------------------------|-----------------------------|
| 1       | R           | C4                  | 15 Hz                 | 57                           | 122               | 90                          | 73.77%                      |
| 2       | L           | C3                  | 16 Hz                 | 49                           | 87                | 49                          | 56.32%                      |
| 3       | R           | C4                  | 19 Hz                 | 72                           | 125               | 43                          | 34.40%                      |
| 4       | R           | C4                  | 11 Hz                 | 38                           | 104               | 91                          | 87.50%                      |
| 5       | R           | C4                  | 11 Hz                 | 64                           | 98                | 82                          | 83.67%                      |
| 6       | L           | C3                  | 9 Hz                  | 62                           | 74                | 21                          | 28.38%                      |
| 7       | L           | C3                  | 15 Hz                 | 57                           | 112               | 104                         | 92.86%                      |
| 8       | L           | C3                  | 11 Hz                 | 68                           | 465               | 333                         | 71.61%                      |
| 9       | R           | C4                  | 11 Hz                 | 37                           | 120               | 55                          | 45.83%                      |
| 10      | L           | C3                  | 17 Hz                 | 66                           | 187               | 185                         | 98.93%                      |
## Supplemental Table II. Summary of Outcome Measures.

| Outcome Measure               | Baseline Score            | Exit Score              | Score Change          | p     |
|-------------------------------|---------------------------|-------------------------|-----------------------|-------|
| Grip Strength (lbs)           | 14.70 (15.80) ± 6.86      | 18.03 (18.30) ± 7.67    | 3.32 (2.70) ± 4.23    | 0.046 |
| Pinch Strength (lbs)          | 1.68 (0.00) ± 2.23        | 4.38 (0.50) ± 5.40      | 2.70 (0.50) ± 4.73    | 0.125 |
| Motricity Index               | 39.8 (37.0) ± 15.5        | 51.9 (51.0) ± 19.9      | 12.1 (12.0) ± 13.4    | 0.027 |
| Motricity Index (Pinch)       | 7.4 (5.5) ± 8.5           | 13.9 (11.0) ± 13.6      | 6.5 (0) ± 10.12       | 0.125 |
| Motricity Index (Elbow)       | 17.9 (19.0) ± 4.8         | 21.2 (25.0) ± 4.8       | 3.3 (5.5) ± 5.5       | 0.13  |
| Motricity Index (Shoulder)    | 13.4 (14.0) ± 3.9         | 15.8 (14.0) ± 3.8       | 2.3 (0.0) ± 2.8       | 0.125 |
| ARAT Total                    | 13.4 (10.1) ± 10.25       | 19.6 (16.0) ± 12.2      | 6.2 (6.0) ± 4.4       | 0.002 |
| ARAT Grasp (Max=18)           | 4.3 (2.75) ± 4.3          | 6.7 (6.0) ± 4.7         | 2.4 (2.0) ± 2.1       | 0.016 |
| ARAT Grip (Max=12)            | 3.6 (3.0) ± 2.3           | 5.4 (4.5) ± 3.0         | 1.9 (1.5) ± 1.6       | 0.004 |
| ARAT Pinch (Max=18)           | 1.5 (0.0) ± 2.8           | 2.3 (0.0) ± 4.2         | 0.9 (0.0) ± 1.5       | 0.250 |
| ARAT Gross (Max=9)            | 4.1 (4.0) ± 1.7           | 5.2 (5.5) ± 1.7         | 1.1 (0.0) ± 1.4       | 0.125 |
| Modified Ashworth Scale       | 1.17 (1.25) ± 0.76        | 1.28 (1.00) ± 0.97      | 0.11 (0.00) ± 0.75    | 0.875 |
| Active Range of Motion        |                           |                         |                       |       |
| Flexion (Digits 2 & 3)        | 66.7 (70.75) ± 11.8       | 72.9 (75.0) ± 20.1      | 6.3 (6.25) ± 10.1     | 0.099 |
| Extension (Digits 2 & 3)      | 49.1 (36.75) ± 21.0       | 47.1 (55.0) ± 31.2      | -2.0 (-9.25) ± 25.4   | 0.819 |
| Flexion (Digits 4 & 5)        | 63.6 (67.5) ± 19.2        | 69.4 (75.0) ± 22.4      | 5.8 (8.75) ± 11.8     | 0.179 |
| Extension (Digits 4 & 5)      | 41.8 (44.75) ± 22.6       | 42.8 (40.0) ± 29.9      | 1.1 (-1.0) ± 21.0     | 0.884 |
| Canadian Occupational Performance Measure (COPM) | | | | |
| Performance                   | 2.10 (2.0) ± 1.08         | 3.66 (3.3) ± 1.68       | 1.56 (1.6) ± 1.70     | 0.022 |
| Satisfaction                  | 1.26 (1.1) ± 0.43         | 2.80 (2.0) ± 2.10       | 1.54 (0.8) ± 1.86     | 0.031 |

a. All measures are reported as mean (median) ± SD
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