Cerebral Contribution to the Execution, But Not Recalibration, of Motor Commands in a Novel Walking Environment

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Abstract

Human movements are flexible as they continuously adapt to changes in the environment. The recalibration of corrective responses to sustained perturbations (e.g., constant force) altering one’s movement contributes to this flexibility. We asked whether the recalibration of corrective actions involve cerebral structures using stroke as a disease model. We characterized changes in muscle activity in stroke survivors and control subjects before, during, and after walking on a split-belt treadmill moving the legs at different speeds. The recalibration of corrective muscle activity was comparable between stroke survivors and control subjects, which was unexpected given the known deficits in feedback responses poststroke. Also, the intact recalibration in stroke survivors contrasted their limited ability to adjust their muscle activity during steady-state split-belt walking. Our results suggest that the recalibration and execution of motor commands are partially dissociable: cerebral lesions interfere with the execution, but not the recalibration, of motor commands on novel movement demands.

Key words: electromyography; feedback; sensorimotor adaptation; split-belt; stroke

Significance Statement

Corrective responses mediated by feedback have been shown to adapt according to task demands. They also reflect updates in the recalibration of the motor system to sustained and predictable changes in the environment. The extent of cortical involvement in this process is unknown. Here we demonstrate that cortical lesions from stroke alter the execution of motor patterns, but not their recalibration. This is important since it suggests that stroke survivors retain the potential to correct movements through error-based protocols, which is an ability that could be exploited for rehabilitation purposes.

Introduction

Humans continuously adapt their movements to changes in the body or environment through corrective responses and adjustment of planned actions. Corrective responses are rapidly triggered on unexpected movement disturbances (Jordan and Rumelhart, 1992; Bhushan and Shadmehr, 1999). Conversely, planned actions are predictive in nature and are updated through sustained perturbations (e.g., constant force) altering one’s movement (Wolpert et al., 1998). Recent work has shown that corrective motor commands also adapt to persistent changes in the environment, such that the subjects perceive the novel situation as the new “normal” (Iturralde and Torres-Oviedo, 2019). However, little is known about...
the neural processes contributing to the recalibration of corrective responses.

It has been suggested that planned and corrective actions share an internal representation of the environmental dynamics (Wagner and Smith, 2008; Maeda et al., 2018); thus, their recalibration could rely on updates to these internal models (Wolpert et al., 1998). If so, the recalibration of corrective responses after sustained exposure to a novel environment is likely dependent on cerebellar structures (Smith and Shadmehr, 2005; Morton and Bastian, 2006), but not on cerebral structures (Reisman et al., 2007; Choi et al., 2009). On the other hand, corrective responses are cerebral dependent, as evidenced by the deficient nature of corrective responses in stroke survivors (i.e., poor muscle coordination, amplitude, or latency; Marigold and Eng, 2006; De Kam et al., 2017, 2018) and impaired task-dependent modulation of this corrective activity (Trumbower et al., 2013; De Kam et al., 2018). Thus, it is plausible that the recalibration of corrective responses is also affected after cerebral lesions, which would imply cerebral-dependent adaptation of corrective actions. Here, we evaluate the involvement of cerebral structures in the recalibration of reactive control through the analysis of corrective muscle activity in individuals with cerebral lesions after stroke.

We characterized stroke-related deficits in muscle activity before, during, and after split-belt walking, which induces robust locomotor adaptation (Reisman et al., 2007). We hypothesized that the execution of motor patterns in a novel walking situation, and the subsequent recalibration of corrective responses would be impaired poststroke. This was based on literature indicating cerebral-related deficits in the modulation of corrective responses (De Kam et al., 2018) and poor muscle coordination poststroke in general (Bowden et al., 2010; Clark et al., 2010; Cheung et al., 2012). Should our hypothesis be supported, our results would suggest that cerebral structures are involved in both the execution and recalibration of corrective actions that result from extended exposure to novel environmental demands.

Materials and Methods

Subjects

We tested 16 stroke survivors in the chronic phase (>6 months) with unilateral supratentorial lesions (i.e., without brainstem or cerebellar lesions; age, 62 ± 9.9 years; 6 females; Table 1) and 16 age- and gender-matched control subjects (age, 61 ± 9.7 years; 6 females). We applied the following inclusion criteria: (1) be able to walk with or without a hand-held device at a self-paced speed for at least 5 min; (2) have no orthopedic or pain conditions interfering with the assessment; (3) have no neurologic conditions except stroke; (4) have no severe cognitive impairments (defined as mini-mental state examination score <24); (5) have no contraindications for performing moderate intensity exercise; and (6) use no medication that interferes with cognitive function. We excluded from data analysis 4 of the 32 participants invited for testing. One stroke participant (P7) was excluded because of severe muscle atrophy and weakness on the sound limb (i.e., non-paretic side), which was present prior to the brain lesion. Another stroke participant (P3) was excluded because of poor muscle recordings due to technical difficulties during testing. One control participant (C1) was excluded because this person failed to follow the testing instructions. Last, we had to remove C7 (i.e., age-matched control of P7) because our regression analyses required equal sample sizes across groups. Namely, including fewer participants in the regression of one group reduces the regressor estimates due to more noise in the averaged data. The study protocol was approved by the Institutional Review Board at the University of Pittsburgh. All study participants gave written informed consent prior to participation.

Experimental setup and protocol

We investigated how participants adapted their kinematic and muscle activation patterns on an instrumented split-belt treadmill (Bertec) with two belts that moved at either the same speed (tied condition) or at different speeds (split condition). We first measured subjects’ overground walking speed using the 6 min walking test (Rikli and Jones, 1998; Kervio et al., 2003), and we then performed the Fugl-Meyer assessment (Fugl-Meyer et al., 1975). Subsequently, subjects walked on the split-belt treadmill. We kept the mean speed across the belts constant in the tied and split conditions. Each subject’s mean belt speed was set to 0.35 m/s below their overground walking speed during the 6 min walking test, yielding a comfortable speed for treadmill walking. The mean belt speed, denoted as medium speed, is reported for each subject in Table 1. In the split condition, the speed of one belt was decreased (slow belt) and the speed on the other belt was increased (fast belt) by 33% of the medium speed to obtain a belt speed ratio of 2:1. Stroke survivors walked with their paretic leg on the slow belt, whereas healthy subjects walked with their nondominant leg on the slow belt. The treadmill protocol consisted of five periods: (1) 50 strides (i.e., time between two subsequent heel-strikes of the same leg) walking at medium speed to familiarize subjects with treadmill walking; (2) a short exposure (10 strides) to the split condition to allow subjects to briefly experience the split condition prior motor adaptation; (3) 150 strides of baseline walking at medium speed to characterize subjects’ baseline gait; (4) 900 strides of adaptation to the split condition, which is a long enough period for locomotor adaptation (Itturralde and Torres-Oviedo, 2019); and (5) 600 strides for afteradaptation at medium speed to measure adaptation effects (i.e., aftereffects) and their decay (Fig. 1A). Subjects had several...
resting breaks during the experiment and some stroke individuals completed fewer strides during adaptation and afteradaptation to prevent fatigue (Table 1 shows the number of strides completed per subject). Participants wore a safety harness, not supporting body weight, attached to a sliding rail in the ceiling to prevent falls. Moreover, subjects could hold on to a handrail in front of the treadmill, but were instructed to do so only if needed.

Data collection

We collected kinetic, kinematic, and electromyogram (EMG) data to characterize individuals’ walking pattern. The ground reaction force aligned with gravity (Fz; sampled at 1000 Hz) was used to identify the instants at which the feet landed (i.e., heel-strike: Fz > 10 N) or were lifted from the ground (i.e., toe-off: Fz < 10 N; Iturralde and Torres-Oviedo, 2019). The positions of the ankles (lateral malleolus) and hips (greater trochanter) were recorded at 100 Hz using a 3D motion analysis system (Vicon Motion Systems). The activity of 15 muscles was recorded bilaterally at 2000 Hz using a Delsys Trigno System (Delsys): gluteus medius, tensor fasciae latae, adductor magnus, hip flexors, rectus femoris, vastus lateralis, vastus medialis, semitendinosus, semimembranosus, biceps femoris, gastrocnemius medialis, gastrocnemius lateralis (LG), soleus, peroneus, and tibialis anterior. EMG signals were high-pass filtered with a 30 Hz fourth-order Butterworth dual-pass filter and subsequently rectified (Merletti and Parker, 2005).

Data analysis

Kinematic parameters

We characterized the adaptation of step-length asymmetry (StepAsym; Eq. 1; Fig. 1C), which is conventionally used to quantify gait changes during split-belt walking (Reisman et al., 2007; Torres-Oviedo et al., 2011). We defined StepAsym as the difference between consecutive steps of the legs in terms of step length, where step length is the distance between the feet (i.e., ankle markers) at heel-strike. In our definition, StepAsym is positive when the step length of the fast leg (i.e., dominant or nonparetic) is larger than the one of the slow leg (nondominant or paretic). We also quantified spatial (StepPosition) and temporal (StepTime) gait features that contribute to StepAsym, since those are differentially affected across stroke survivors, and they exhibit distinct adaptation patterns in unimpaired adults during split-belt walking (Finley et al., 2015). Finally, StepVelocity was defined as the difference between the legs in terms of velocity of the foot with respect to the body when in contact with the ground. All parameters were expressed in units of distance, and they were normalized to the sum of left and right step lengths to account for differences in step sizes across subjects (Sombric et al., 2017).

EMG parameters

We characterized the modulation of muscle activity across the different walking conditions using the average activity of each muscle for fixed phases of the gait cycle (Fig. 1C). Specifically, we divided the gait cycle into the following four phases: first double support (DS; between ipsilateral heel-strike and contralateral toe-off), single stance (SINGLE; from contralateral toe-off to contralateral heel-strike); second DS (between contralateral heel-strike and ipsilateral toe-off) and swing (SWING; between ipsilateral toe-off and ipsilateral heel-strike). We further divided each of these phases to achieve better temporal resolution. Specifically, both DS phases were divided in two equal subphases, and the SINGLE and SWING

| Subject | Age (years) | Gender | Affected side | Lesion location | Fugl-Meyer score | Medium walking speed | Adapt strides | Post strides |
|---------|-------------|--------|---------------|----------------|-----------------|---------------------|---------------|-------------|
| P1      | 43          | Female | R             | Left MCA and basal ganglia | 33              | 1.13                | 907           | 605         |
| P2      | 64          | Female | R             | Left MCA and ACA, temporal lobe, basal ganglia | 26              | 0.81                | 867           | 642         |
| P3      | 64          | Female | R             | Left MCA, frontal, parietal lobe and basal ganglia | 29              | 0.60                | 616           | 308         |
| P4      | 58          | Female | R             | Left medial, frontal and parietal area’s | 21              | 0.45                | 901           | 624         |
| P5      | 56          | Female | L             | Right parietal posterior and temporal lobes | 31              | 0.94                | 941           | 615         |
| P6      | 64          | Male   | L             | Right MCA | 31              | 0.34                | 452           | 300         |
| P7      | 78          | Male   | R             | Right MCA | 486             | 217                 |               |             |
| P8      | 55          | Female | L             | Right MCA | 23              | 0.87                | 903           | 602         |
| P9      | 66          | Male   | R             | Left MCA, frontal, temporal and parietal lobes | 30              | 0.77                | 605           | 599         |
| P10     | 60          | Female | R             | Left frontal | 26              | 0.9                 | 908           | 600         |
| P11     | 77          | Male   | R             | Thalamus | 30              | 0.35                | 590           | 601         |
| P12     | 59          | Male   | R             | Left MCA | 32              | 0.7                 | 905           | 600         |
| P13     | 52          | Male   | R             | Left MCA | 32              | 0.96                | 903           | 603         |
| P14     | 66          | Male   | L             | Right frontal superior, parietal and posterior area steady state | 29              | 0.76                | 909           | 602         |
| P15     | 75          | Male   | R             | Left periventricular, temporal and basal ganglia | 32              | 0.94                | 913           | 552         |
| P16     | 49          | Male   | R             | Frontotemporal parietal | 33              | 0.71                | 931           | 303         |

ACA, anterior cerebral artery; MCA, middle cerebral artery. Clinical characteristics of stroke survivors. Bold type indicates subjects who are included in the speed-matched analysis. Italic type indicates subjects who are included in the asymmetry-matched analysis. Bold-italic type indicates subjects who are included in both speed-matched analysis and asymmetry-matched analysis.
phases were subdivided in four equal subphases. Muscle activity amplitude was averaged in time for each of these subintervals for every stride and muscle, resulting in 180 muscle activity variables per leg per stride cycle: 12 sub-interval/C2 15 muscles.

EMG activity for each muscle was linearly scaled to baseline walking (last 40 strides), such that a value of 0 corresponded to the average of the interval with the lowest average activity and 1 corresponded to the average of the interval with the highest average activity (Iturralde and Torres-Oviedo, 2019). This normalization enabled us to aggregate the EMG activity across subjects to perform group analyses. Of note, we excluded from analysis the activity of soleus from one stroke survivor because technical difficulties during data collection.

Epochs of interest

Kinematic and EMG parameters were used to characterize subjects’ behavior at the beginning (“early”) and at the end (“late”) of each experimental condition. Specifically, the epochs of interest included the following: late baseline walking, early and late adaptation, and early afteradaptation. The early epochs were characterized by the median of the initial five strides and late epochs by the median of the last 40 strides of the condition of interest. We chose medians across strides, rather than means to minimize the impact of outlier values. In all cases, we excluded the very first and very last stride of each condition to avoid artifacts from starting and stopping the treadmill. Subsequently, we subtracted the late baseline behavior from all epochs of interest. This allowed us to identify group differences in subjects’ modulation of kinematic and EMG parameters beyond those due to distinct baseline biases. Moreover, we computed the differences between EMG activity early afteradaptation versus late adaptation to quantify changes in EMG activity on sudden removal of the perturbation.

Sensorimotor recalibration of corrective muscle responses

We studied the structure (i.e., activity across multiple muscles) of corrective motor responses on sudden changes in the walking environment (Fig. 2), since this reflects the extent of sensorimotor recalibration (Iturralde and Torres-Oviedo, 2019). We defined corrective responses as the rapid changes in motor output (ΔEMG) immediately after a transition in the walking environment. Corrective responses were quantified as the difference in muscle activity immediately after an environmental transition (EMGafter) compared with the muscle activity before the transition (EMGbefore; Fig. 2A). Thus, corrective response (ΔEMG) = EMGafter − EMGbefore. Since we had multiple strides before and after a transition, we used the median EMG activity across either 40 or 5 strides to quantify EMGbefore and EMGafter given a transition, respectively.

Corrective responses, ΔEMG, were labeled according to the environmental transition and split-belt environment that was inducing them. More specifically, ΔEMG on referred to corrective responses when the split environment was introduced (“on” transition), and ΔEMG off referred to those when the split environment was removed (“off” transition). Also, corrective responses were labeled “(+)” or “(−)” to indicate the specific split environment that was
Figure 2. Environment-based and adaptive contributions to corrective responses. A, Schematic representation of corrective responses on the introduction split-belt perturbation. The split environment is arbitrarily defined as + if the paretic leg is on the slow belt and the nonparetic leg is on the fast belt, whereas it is defined as – if the paretic leg is on the fast belt and the nonparetic leg is on the slow belt. Changes in muscle activity (i.e., corrective response) on the introduction of the + or – environment are color coded as blue (decreased activity), red (increased activity), and white (no change in activity). B, In the case of an environment-based corrective response changes in muscle activity perturbation removal (∆EMG_{off}) are opposite to those on perturbation introduction. C, In the case of an adaptive corrective response, the split environment is perceived as the new normal. Consequently, removal of the split environment will be experienced as a perturbation in the opposite direction. Thus, the structure of the corrective response will resemble the one observed on introduction of the – environment. D, Regression equation used to quantify the structure of corrective response ∆EMG_{off}. In this equation, β_{adapt} quantifies the similarity of ∆EMG_{off} to the adaptation-based response and β_{no-adapt} quantifies the similarity of ∆EMG_{off} to the environment-based response.

We were specifically interested in the structure of corrective responses postadaptation because this structure indicates the extent to which subjects recalibrate their motor system (Iturralde and Torres-Oviedo, 2019). Namely, the structure of these corrective responses is determined by both changes in the environment and changes in the motor systems’ adaptive state. We discerned the environment-based and adaptive-based contributions to corrective responses afteradaptation (∆EMG_{off}) with a regression model (∆EMG_{off} = adaptive-based + environment-based + ε). In the case of an environment-based response, the corrective pattern ∆EMG_{on} on introducing the ‘+’ split environment is simply disengaged once this environment is removed (i.e., both belts moving at the same speed; Iturralde and Torres-Oviedo, 2019). Thus, in this case the structure of corrective responses afteradaptation ∆EMG_{off} (i.e., when the split ‘+’ environment is turned off) resembles the numerical opposite of ∆EMG_{on} (ΔEMG_{off} = −ΔEMG_{on}; Fig. 2B). Conversely, adaptive-based responses are observed if subjects perceive the split-belt environment (+) as the “new normal.” Consequently, removing the (+) environment is processed as a perturbation in the opposite direction as the one originally experienced [i.e., it would be equivalent to introduction of the (−) environment; Fig. 2A]. Thus, in the case of adaptive corrective responses, the structure of ∆EMG_{off} resembles corrective responses to transitioning into the opposite (−) split-belt environment (∆EMG_{off} = ∆EMG_{on}; Fig. 2C). Note that the corrective responses afteradaptation (∆EMG_{off}) exhibit features of both environment-based and adaptive-based responses. Thus, we used a regression analysis to determine the extent to which the structure of corrective responses afteradaptation was environment-based or adaptive-based (Fig. 2D), as follows:

$$\Delta EMG_{\text{off}} = \beta_{\text{no-adapt}} \Delta EMG_{\text{on}} + \beta_{\text{adapt}} \Delta EMG_{\text{on}} + \varepsilon$$

In the regression equation, the parameters $\beta_{\text{no-adapt}}$ and $\beta_{\text{adapt}}$ are respectively interpreted as the extent to which the structure of corrective responses indicates transitions in the environment (i.e., environment based) or the adaptation of subjects’ motor system (i.e., adaptive based). Note that every vector is divided by its norm (i.e., $\Delta EMG_{\text{off}} = \Delta EMG_{\text{off}} / \| \Delta EMG_{\text{off}} \|$). This was done because we were interested in identifying stroke-related deficits in the structure, rather than the magnitude of corrective responses, which is known to be different (De Kam et al., 2017). For example, we find that the amplitude of
corrective responses, $\Delta EMG_{on(\cdot)}$, for each leg was smaller for the stroke group ($\Delta EMG_{on(\cdot)} = 2.6$ and 2.2) than the control group (3.3 and 3.7).

Note that $\Delta EMG_{on(\cdot)}$ was not directly measured to avoid exposing subjects to multiple environmental transitions prior to the adaptation period. Instead, we inferred these responses by exploiting the symmetry of the transition between the two legs. The only difference between the + and − environments is which leg increases speed and which leg decreases it. We used this similarity to infer the (not recorded) corrective responses ($\Delta EMG_{on(\cdot)}$) of each leg to transitioning into the − environment from the (measured) corrective responses ($\Delta EMG_{on(\cdot)}$) to transitioning to the + environment. In other words, we assumed that the (not recorded) nondominant leg’s responses to the “on (−)’’ transition would be similar to the (recorded) dominant leg’s responses to the “on (+)’’ transition, and vice versa. We are aware that this assumption might not be valid for some afterstroke individuals, given their inherent motor asymmetry. Thus, group differences in $\beta_{adapt}$ values, which are estimated using the not recorded $\Delta EMG_{on(\cdot)}$ in our regression analysis, might be due to the experimental limitation of our study. To address this possibility, we performed a post hoc analysis to compare the regression coefficients between a subset of patients and control subjects ($n = 7$ on each subgroup) that had similar asymmetry in their EMG activity during baseline walking ($p = 0.1$). The baseline asymmetry in EMG activity across the legs was quantified in each subject by first computing a 180-dimension vector (15 muscles × 12 gait cycle phases) of the baseline muscle activity for each leg and then calculating the cosine between those baseline vectors for the legs of each individual.

Structure of muscle activity patterns in a novel walking environment

We characterized changes in the structure of steady-state muscle activity from baseline walking to late adaptation ($\Delta EMG_{SS} = EMG_{late adaptation} − EMG_{baseline}$). This was defined as the pattern of activity across all muscles and all gait cycle intervals (15 muscles × 12 intervals = 180 data points for a given epoch). The $\Delta EMG_{SS}$ 180-dimensional vector for each subject was used to assess structural differences between stroke survivors and control subjects. We specifically computed a cosine between the $\Delta EMG_{SS}$ for each individual and a “reference pattern” $\Delta EMG_{SS}$, which was defined as the median $\Delta EMG_{SS}$ of the control group. This reference pattern for $\Delta EMG_{SS}$ was calculated as the group median of all control subjects when computing the similarity metric for each leg of the stroke survivors, whereas for individual control subjects we excluded the subjects’ own data to compute the reference vector. A cosine closer to 1 indicates that the subject-specific and “reference” vectors are more aligned, and therefore, the structures of the muscle patterns that they represent are similar.

Statistical analyses

Modulation of muscle activity within groups

Modulation of muscle activity was first evaluated for each group individually. Specifically, we compared muscle activity between the epochs of interest using a Wilcoxon signed-rank test (nonparametric equivalent of paired $t$ test) for each individual muscle and for each gait cycle phase, resulting in 360 comparisons per epoch (12 intervals × 15 muscles × 2 legs). We subsequently corrected the significance threshold for each epoch using a Benjamini–Hochberg procedure (Benjamini and Hochberg, 1995) to indicate significant changes in our figures, but all data in both groups was used in the structural analyses.

Structure of muscle activity patterns during steady-state walking

We used a Wilcoxon rank sum test to compare the groups on their $\Delta EMG_{SS}$ for each leg during late adaptation in the split-belt condition. We specifically compared the group’s similarity in $\Delta EMG_{SS}$ to the reference pattern obtained with the cosine analysis.

Sensorimotor recalibration of corrective muscle responses

We compared the regressor coefficients $\beta_{no-adapt}$ and $\beta_{adapt}$ for each group to determine whether stroke survivors and control subjects differed in the adaptation of corrective responses. Since the regressor estimates of $\beta_{no-adapt}$ and $\beta_{adapt}$ in a regression model are not independent, between-group comparisons were performed in the 2D space covered by $\beta_{no-adapt}$ and $\beta_{adapt}$. The differences between the groups were compared using a $\chi^2$ distribution, which could be considered as a high-dimensional $t$ test (Härdle and Simar, 2007).

Correlation analyses

We asked whether individual subjects’ adaptation of muscle activity was related to the severity of motor impairment (i.e., Fugl-Meyer score). To this end, we performed Spearman correlations between (1) the Fugl-Meyer score and (2) outcome measures that reflected sensorimotor recalibration (i.e., $\beta_{adapt}$ and $\beta_{no-adapt}$) and the similarity metric comparing the structure of muscle activity during late adaptation in the split-belt condition for each individual versus a reference $\Delta EMG_{SS}$.

Modulation of kinematic parameters

We compared stroke survivors and control subjects in how they modulated kinematic parameters. To this end, we performed a repeated-measures analysis of variance for each kinematic outcome (StepAsym, StepPosition, StepTime, and StepVelocity) with GROUP (stroke vs control subjects), EPOCH (early adaptation, late adaptation, and early postadaptation), and the interaction between both variables as predictors. Note that we did this analysis with unbiased data (i.e., baseline subtracted) because we were interested in differences in modulation across groups, beyond their baseline biases. In case of a significant GROUP or GROUP × EPOCH interaction effect, we performed between-group comparisons for each epoch using Bonferroni-corrected independent $t$ tests (adjusted $\alpha = 0.017$).
Speed-matched post hoc analysis
We found that stroke survivors walked slower than control subjects during the experiment (averaged medium speed = 0.78 ± 0.24 vs 1.07 ± 0.12 m/s; rank-sum test, p < 0.01), which could confound between-group differences in muscle activity. Thus, we repeated our analyses with only the 10 fastest participants in the stroke group and the 10 slowest control subjects to determine whether structural differences between our groups were due to walking speed, rather than brain lesion. Walking speed was not significantly different for these speed-matched subgroups (0.88 ± 0.18 vs 1.0 ± 0.15 m/s; rank sum test, p = 0.10). Importantly, the selection of the fastest stroke survivors did not result in a selection of patients with less severe motor impairments (Fugl-Meyer score = 29.5). We did not find between-group differences in muscle activity at steady state was similar to the one of control subjects, whereas the paretic leg was not (Fig. 3A). Differences in the structure of muscle activity modulation between the groups can be appreciated in Figure 3B. Specifically, similarity metric ΔEMGSS was lower in the paretic leg compared with control subjects (Fig. 3A; p = 0.001), and between-group differences were trending (p = 0.057) when comparing the nonparetic leg activity to that of control subjects. These between-group differences were not observed when patients and control subjects walked at similar speeds (median ± interquartile range in control subjects vs stroke survivors for the nonparetic leg: 0.58 ± 0.26 vs 0.51 ± 0.22, p = 0.47; paretic leg: 0.39 ± 0.13 vs 0.28 ± 0.18, p = 0.1). Interestingly, a more atypical structure in muscle activity modulation in the paretic leg was associated with poorer voluntary leg motor control as measured by the Fugl-Meyer scale (p = 0.59, p = 0.028, Fig. 3C), but not in the nonparetic leg (p = −0.29, p = 0.32; data not shown). In conclusion, the structure of muscle activity at steady state was different between patients and control subjects, and individuals with more atypical paretic activity were those with lower voluntary function.

Results
Cerebral lesions interfered with the structure of muscle activity in a novel walking environment
We computed a similarity metric, ΔEMGSS, which indicated the similarity between the structure of an individual’s muscle activity modulation in steady-state walking relative to the average pattern in control subjects (reference pattern). We found that the nonparetic leg activity at steady state was similar to the one of control subjects, whereas the paretic leg was not (Fig. 3A). Differences in the structure of muscle activity modulation between the groups can be appreciated in Figure 3B. Specifically, similarity metric ΔEMGSS was lower in the paretic leg compared with control subjects (Fig. 3A; p = 0.001), and between-group differences were trending (p = 0.057) when comparing the nonparetic leg activity to that of control subjects. These between-group differences were not observed when patients and control subjects walked at similar speeds (median ± interquartile range in control subjects vs stroke survivors for the nonparetic leg: 0.58 ± 0.26 vs 0.51 ± 0.22, p = 0.47; paretic leg: 0.39 ± 0.13 vs 0.28 ± 0.18, p = 0.1). Interestingly, a more atypical structure in muscle activity modulation in the paretic leg was associated with poorer voluntary leg motor control as measured by the Fugl-Meyer scale (p = 0.59, p = 0.028, Fig. 3C), but not in the nonparetic leg (p = −0.29, p = 0.32; data not shown). In conclusion, the structure of muscle activity at steady state was different between patients and control subjects, and individuals with more atypical paretic activity were those with lower voluntary function.

Sensorimotor recalibration of corrective responses was intact after cerebral lesion
The structure of corrective responses for each group indicated that on average both groups recalibrated their gait similarly. This is qualitatively indicated by the “checker boards” illustrated in Figure 4. Notice that in both groups the observed corrective responses postadaptation (Fig. 4C) look more similar to those predicted by the adaptive (Fig. 4B) than the environment-based modulation (Fig. 4A). The environment-based and adaptive-based contributions to corrective responses postadaptation were quantified with a regression model, which reproduced the data well (Fig. 5, left panels). We observed that the regression coefficient \( \beta_{\text{adapt}} \) was greater than \( \beta_{\text{no-adapt}} \) in both groups for the leg that walked slow (i.e., nondominant leg in control subjects: CI for \( \beta_{\text{adapt}} = 0.68–0.85 \) vs CI for \( \beta_{\text{no-adapt}} = 0.18–0.35 \); paretic leg in stroke: CI for \( \beta_{\text{adapt}} = 0.55–0.77 \) vs CI for \( \beta_{\text{no-adapt}} = 0.10–0.32 \)) and the leg that walked fast (i.e., dominant leg in control subjects: CI for \( \beta_{\text{adapt}} = 0.73–0.89 \) vs CI for \( \beta_{\text{no-adapt}} = 0.09–0.25 \); nonparetic leg in stroke: CI for \( \beta_{\text{adapt}} = 0.54–0.71 \) vs CI for \( \beta_{\text{no-adapt}} = 0.46–0.62 \)). These coefficients were not different between groups when estimated from the averaged paretic leg activity across stroke survivors vs that of the nondominant leg across control subjects (\( \chi^2 = 3.2, p = 0.20 \)), indicating that averaged responses in the slow leg were adapted to the same extent in stroke survivors and control subjects. Conversely, we found between-group differences when comparing the coefficients of the averaged nonparetic activity in the stroke group versus that of the dominant leg in the control group (\( \chi^2 = 48.9, p = 2.4 \times 10^{-11} \); Fig. 5A, bottom). Thus, we observed between-group differences in the regression coefficients for the nonparetic, but not the paretic, compared with control legs.

As a post hoc analysis, we considered the possibility that these group differences in the nonparetic side could arise from our estimation of the adaptive-based modulation (ΔEMGon–(−)). Notably, this muscle activity was not recorded but it was inferred from the muscle activity of the other leg, assuming symmetry of corrective responses across legs. Given that stroke survivors exhibit asymmetric motor patterns, the ΔEMGon+(+1) of the paretic leg may not be a good estimate for the ΔEMGon(−) of the nonparetic leg, thereby leading to underestimation of \( \beta_{\text{adapt}} \) in this leg. Thus, we performed a subgroup analysis in which stroke survivors and control subjects were matched for symmetry in their muscle activity during baseline walking. We did not find between-group differences for either leg of the stroke group compared with the control subjects when asymmetry in baseline muscle activity was matched between the groups (Fig. 5B; paretic vs nondominant control leg: \( \chi^2 = 4.1, p = 0.13 \); nonparetic vs dominant control legs, \( \chi^2 = 2.5, p = 0.29 \)). In conclusion, the observed structure of corrective responses postadaptation were more similar to the one predicted by adaptive, rather than environment-based, modulation in patients with cerebral lesions and control subjects.

While recalibration of corrective responses poststroke did not differ from that of control subjects at the group level when asymmetries were accounted for, we considered the possibility that some individuals would exhibit less recalibration compared with others. Consistently, Figure 6A shows a wide range of \( \beta_{\text{adapt}} \) and \( \beta_{\text{non-adapt}} \) regression values at the individual level. Also, note that the regression model had smaller \( R^2 \) values when applied to each subject’s corrective responses postadaptation (control subjects’ nondominant leg: \( R^2 = 0.38 \pm 0.18 \); control subjects’ dominant leg: \( R^2 = 0.42 \pm 0.21 \); stroke survivors’ nondominant leg: \( R^2 = 0.66 \pm 0.25 \); stroke survivors’ dominant leg: \( R^2 = 0.48 \pm 0.17 \)).
subjects’ dominant leg: 34 ± 0.17; paretic leg: 0.18 ± 18; nonparetic leg: 0.18 ± 18) than to the corrective response of the group (reported in previous section). However, the regression model was significant in all individuals, except for one stroke survivor (p = 0.19). In sum, we find large ranges of regression coefficients in control and poststroke individuals.

We further asked whether stroke survivors would exhibit less recalibration if they had more severe leg motor impairments (i.e., Fugl-Meyer scale). Thus, we computed the Spearman correlation between individual subjects’ regression coefficients and their leg motor score (Fig. 6B). We found that βadapt of neither the paretic nor nonparetic legs was correlated to the Fugl-Meyer score (paretic: ρ = 0.34, p = 0.23; nonparetic: ρ = 0.23, p = 0.43). On the other hand, motor function measured with the Fugl-Meyer score was associated with βno-adapt for the paretic leg and not the βno-adapt for the nonparetic leg (paretic leg: ρ = 0.60, p = 0.024; nonparetic leg: p = 0.02, p = 0.94). However, this correlation was driven by the individual with the largest negative βno-adapt value since the correlation was no longer significant when this subject was excluded (ρ = 0.37, p = 0.22). As such, we are cautious about interpreting this result as a positive association between environment-based corrective response and leg motor scores. Together, our correlation analyses indicate that recalibration of corrective responses is not associated with the quality of voluntary motor control.

**Stroke-related deficits in muscle coordination are not reflected in asymmetry parameters**

While stroke survivors exhibited deficits in the execution of updated motor commands during steady-state split-belt walking (i.e., ΔEMGSS), we observed no differences between the groups in the modulation of asymmetry parameters (i.e., stepAsym, stepPosition, stepTime, and stepVelocity; Fig. 7). Specifically, we observed no main effects of GROUP or GROUP × EPOCH interaction effects for the interlimb kinematic parameters (p > 0.05). Comparable results were obtained in our speed-matched analysis. Thus, interlimb kinematic parameters are less sensitive to stroke-related deficits in locomotor adaptation than our outcome measures for muscle coordination.

**Discussion**

We studied the involvement of cerebral structures in the sensorimotor recalibration of gait using stroke as a clinical model. We found that on average stroke survivors
exhibit similar recalibration of corrective responses in the paretic leg relative to control subjects, which was surprising given the known deficits in paretic responses post-stroke. On the other hand, we found that cerebral lesions affected the muscle activity of paretic legs in the steady state of split-belt walking. Thus, we find an interesting dissociation between execution and recalibration of corrective actions: the execution of motor patterns on novel demands is cerebral dependent, but the recalibration is not. These results do not support our original hypothesis that execution and recalibration of corrective responses would be a cerebral-mediated process. Our findings suggest as though this latter process might only depend on other structures such as the cerebellum.

**Sensorimotor recalibration of corrective responses after cerebral lesions**

Our results suggest that while corrective responses are affected by lesions to cerebral structures (Marigold and Eng, 2006; De Kam et al., 2017, 2018) and corticospinal tract (Christensen et al., 1999, 2001), the recalibration of this corrective activity is not strongly mediated by these neural structures. More specifically, we found...
that the changes in corrective responses postadaptation are similar between individuals with cerebral lesions and control subjects. This is clearly evident in the paretic leg, but not in the nonparetic leg, which exhibited lower adaptive-based changes (i.e., lower $\beta_{\text{adapt}}$). We speculate that $\beta_{\text{adapt}}$ was lower in patients than in control subjects because it was underestimated due to asymmetry in corrective responses poststroke (Marigold and Eng, 2006; De Kam et al., 2018), rather than because poor recalibration in the nonparetic leg. However, future studies recording $\Delta \text{EMG}_{\text{on}(-)}$ are needed to determine whether motor asymmetry is a factor reducing the recalibration of nonparetic corrective responses.

Discrepancies between the paretic and nonparetic extent of adaptive-based changes may reflect leg-specific recalibration. This is supported by the independent recalibration of the legs in hybrid walking (i.e., one leg moving forward faster than the other leg moving backward; Choi and Bastian, 2007). However, leg-specific adaptation in hybrid walking may result from the peculiar nature of this task and may, therefore, not apply to other locomotor adaptation paradigms. In fact, more recent studies have demonstrated interlimb transfer of adapted motor patterns during conventional split-belt walking (Krishnan et al., 2017, 2018), which argues against leg-specific recalibration. Thus, we believe that recalibration of corrective responses is not affected after cerebral lesions, which is also supported by a lack of association between individual stroke survivors’ motor impairments (i.e., Fugl-Meyer scores) and the amount of adaptive-based modulation of corrective responses. Of note, we found an association between $\beta_{\text{no-adapt}}$ and Fugl-Meyer scores driven by stroke survivors with negative $\beta_{\text{no-adapt}}$ values. These negative $\beta_{\text{no-adapt}}$ values, also observed in control subjects (Iturralde and Torres-Oviedo, 2019), may reflect a startle-like (Oude Nijhuis et al., 2010) generic response to an environmental transition, regardless of its direction. Together, our results suggest that, while corrective responses are affected by cerebral lesions (De Kam et al., 2018), their recalibration is not.

Cerebral lesions affect the execution of updated motor commands in a new walking environment

We found that stroke survivors exhibited impaired modulation of steady-state muscle activity, particularly in their paretic leg. Interestingly, aberrant patterns of muscle activity did not impact the modulation of kinematic asymmetry parameters (Reisman et al., 2007), which may indicate that muscle activity is more sensitive to stroke-related deficits in motor output than parameters quantifying kinematic asymmetries. Previous studies reported that long-term adaptation (i.e., changes in spatiotemporal...
parameters from baseline walking to late adaptation) is impaired after hemispherectomy (Choi et al., 2009), but not after focal hemisphere lesions due to stroke (Reisman et al., 2007). These observations further suggest that changes in interlimb spatiotemporal parameters become noticeable with more pronounced deficits, whereas more subtle deficits can be detected by analysis of muscle activity. The impaired modulation of muscle activity at steady state in the split-belt walking condition may reflect poor selectivity in the activation of muscles poststroke. Notably, intact individuals upregulated proximal muscles (i.e., quadriceps during early stance and hamstrings during late swing) without increasing distal muscle activity during steady-state split-belt walking. This pattern of modulation was diminished in stroke survivors (Fig. 3), particularly those with poorer leg motor selectivity. This observation is consistent with previous reports of missing selectivity in the activation of proximal and distal muscles in the gait of stroke survivors (Clark et al., 2010). Thus, impaired motor selectivity probably contributes to the aberrant patterns of muscle activity during steady-state split-belt walking. In addition, stroke survivors’ atypical steady-state behavior may also be influenced by their perceptual deficits. Notably, individuals poststroke have difficulty assessing their step-length asymmetry (Wutzke et al., 2015), which could contribute to the adaptation of movements (Hoogkamer et al., 2015a). Alternatively, the atypical muscle activity patterns in stroke survivors may have resulted from a lower walking speed. Indeed, steady-state muscle activity became more similar across groups in our speed-matched analysis. Notably, between-group differences in our similarity metric were still substantial after controlling for speed (0.39 ± 0.13 vs 0.28 ± 0.18), but this difference was no longer statistically significant. Therefore, we cannot completely rule out walking speed as a confounding factor influencing the distinct motor patterns at steady state between patients and control subjects. Last, the lack of modulation of steady-state muscle activity was presumably not the result of muscle atrophy, given that muscle groups that lacked modulation in the steady state (e.g., knee extensors) were highly modulated during corrective responses (Fig. 1D). We speculate that steady-state muscle activity depends on neural circuits involved in voluntary motor control, whereas this is not the case for corrective responses (De Kam et al., 2018). Together, our results suggest that stroke survivors exhibit impaired execution of updated motor commands in the steady state of split-belt walking, most likely due to their impaired motor function.

**Partial dissociation between recalibration and execution of updated motor commands**

We found that stroke survivors exhibited intact recalibration of corrective responses, but impaired muscle patterns at steady-state split-belt walking, suggesting partial dissociation between motor performance in the
altered environment and postadaptation behavior. This finding is consistent with previous work demonstrating that the extent to which subjects adapt their movements during split-belt walking does not predict their aftereffects (Sombric et al., 2019). Partial dissociation between steady-state and postadaptation behavior is further supported by the findings that aftereffects are not sensitive to manipulation of steady-state behavior through visual feedback (Wu et al., 2014; Long et al., 2016). Together, our findings suggest that steady-state and postadaptation behaviors are partially independent, and possibly mediated through distinct neural processes.

We found that postadaptation muscle activity was indicative of sensorimotor recalibration of corrective responses also observed in previous studies (Maeda et al., 2018; Iturralde and Torres-Oviedo, 2019). Since recalibration has also been observed in feedforward motor commands on perturbation removal (Tseng et al., 2007; Taylor and Ivry, 2014), our results provide further evidence for shared internal models for generating corrective responses and feedforward motor commands (Wagner and Smith, 2008; Yousif and Diedrichsen, 2012; Cluff and Scott, 2013; Maeda et al., 2018). It has been shown that the cerebellum is involved in feedforward adaptation and learning of internal models. (Martin et al., 1996; Smith and Shadmehr, 2005; Morton and Bastian, 2006). In particular, sensorimotor recalibration in locomotion depends on the intermediate cerebellum (Darmohray et al., 2019), and small focal lesions may not affect it (Hoogkamer et al., 2015b). In addition, the cerebellum may also be involved in the adaptation of corrective responses. This is supported by the cerebellar dependency on timely recruitment (Herzfeld et al., 2014) and the appropriate magnitude of feedback responses to predictable perturbations (Jacobs and Horak, 2007). Together, our results are consistent with the idea that, corrective responses depend on spinal cord and brainstem circuits for their execution (Bolton, 2015; Weiler et al., 2019) and on the cerebellum for their

Figure 7. Modulation of kinematic parameters. A, Group-averaged time courses for StepPosition, StepTime, StepVelocity, and StepAsym. Note that individual subjects’ baseline biases were subtracted to allow for comparison of modulation of parameters regardless of differences in baseline asymmetry. Shaded areas represent SEs for each group. For visual purposes, data were smoothed using a running average (median) of 10 strides. Rectangles represent the epochs of interest. B, Interlimb kinematic parameters for each group during baseline. C, Between-group comparisons for kinematic parameters over the epochs of interest. We found no significant differences between the groups in any of the parameters. StepAsym (GROUP: F(1,28) = 3.48, p = 0.07; GROUP X EPOCH: F(2,56) = 1.16, p = 0.31), stepPosition (GROUP: F(1,28) = 2.14, p = 0.16; GROUP X EPOCH: F(2,56) = 2.33, p = 0.13), stepTime (GROUP: F(1,28) = 3.17, p = 0.09; GROUP X EPOCH: F(2,56) = 0.63, p = 0.50), and stepVelocity (GROUP: F(1,28) = 1.41, p = 0.25; GROUP X EPOCH: F(2,56) = 1.11, p = 0.34).
adaptation, which would explain why our participants with cerebral lesions showed intact recalibration of corrective responses.

Our observation of stroke-related impairments in steady-state movement execution suggest that these processes are cerebral dependent, perhaps through connections between cerebral and cerebellar structures (Kelly and Strick, 2003; Hoshi et al., 2005). Moreover, intact motor pathways for voluntary motor control (e.g., corticospinal tract) are most likely involved in the execution of steady-state motor commands (Schweighofer et al., 2018), given our finding that individuals with poorer voluntary motor control also exhibited a more atypical structure of their steady-state muscle activity. Such associations were not found for the execution of corrective responses (De Kam et al., 2018), suggesting that the execution of corrective responses uses a different circuitry, most likely at the level of the brainstem (Jacobs and Horak, 2007; Bolton, 2015). Together, our results are consistent with the idea that corrective and planned actions share an internal model, which relies on cerebellar structures for their adaptation and on cerebral structures for their execution.

Clinical implications

Our detailed characterization of muscle activity modulation during and after split-belt walking allows for the identification of muscle activity that could potentially be targeted by split-belt treadmill training. Our results support previous findings reporting movement aftereffects in stroke survivors comparable to those of control subjects (Reisman et al., 2007, 2013; Lauzière et al., 2016; Lewek et al., 2018). We further show that the extent of sensorimotor recalibration underlying these aftereffects vary greatly across poststroke individuals. We speculate that individual differences in sensorimotor recalibration may explain why some stroke survivors improve their gait symmetry in response to repeated split-belt treadmill training while others do not (Reisman et al., 2013; Betschart et al., 2018; Lewek et al., 2018). If so, it may be possible to identify patients that will benefit from split-belt training within just a single session. Future studies are needed to determine whether individuals' recalibration of corrective responses can predict their response to repeated training.

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