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Corticomuscular Adaptation to Mechanical Perturbations in a Seated Locomotor Task

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CORTICOMUSCULAR ADAPTATION TO MECHANICAL PERTURBATIONS IN A SEATED LOCOMOTOR TASK

by

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A dissertation submitted in partial fulfilment of the requirements for the degree of Doctor of Philosophy in Mechanical Engineering in the Department of Mechanical and Aerospace Engineering in the College of Engineering and Computer Science at the University of Central Florida
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Major Professor: Helen J. Huang
Cortical control during walking is most pronounced when the person is perturbed. Although seated locomotor tasks such as cycling or recumbent stepping improve walking performance, the electrocortical correlates of perturbed seated tasks have not been studied in detail. The primary purpose of this research was to quantify cortical and muscular responses to mechanical perturbations during recumbent stepping. We also aimed to quantify possible differences between young and older adults’ responses to perturbed stepping. A secondary aim of this research was to determine the accuracy of electroencephalography (EEG) source imaging to interpret the electrocortical findings adequately. We hypothesized that both young and older adults would adapt to the perturbations by reducing their movement errors and reducing the anterior cingulate electrocortical activity. We also hypothesized that older adults would co-contract their agonist and antagonist muscles more than young adults in response to perturbations. Such stronger co-contraction would indicate older adults would have weaker corticomescular connectivity in response to perturbations than young adults.

Seventeen young adults and eleven older adults completed four perturbed arms and leg stepping tasks. We perturbed the stepping with brief 200ms increased movement resistance using a controllable servomotor on our recumbent stepper. We asked subjects to step smoothly, use both arms and legs and follow the visual pacing cue set at 60 steps per minute. We recorded brain activity with high-density EEG with 128 electrodes, muscular activity with 16 electromyography (EMG) sensors, and stepping kinematics using the servomotor’s encoder. We quantified temporal and spatial motor errors from the stepping kinematics data. We used a novel post-processing approach to reject noise from EEG and estimated the electrocortical sources using independent component analysis and the current dipole estimation technique. We then performed a series of time-frequency analyses on the group EEG source data. We quantified EMG co-contraction for each of the perturbed and recovery steps. Finally, we used direct Directed Transfer Function to determine the corticomuscular
connectivity time-locked to young and older adults’ perturbations.

Quantifying the accuracy of source estimation showed that recording the three-dimensional EEG electrode locations would provide accurate source estimation up to a single Brodmann area. We also found that recording the precise location of the fiducials, i.e., the anatomical landmarks used to place the EEG electrodes, is critical for a reliable source estimation process. Motor errors did not show a reduction of errors with more perturbation experience for both young and older adults. Young adults showed significant theta-band (3-8 Hz) electrocortical activity locked to the perturbations at the anterior cingulate cortex, supplementary motor areas, and posterior parietal cortex. These locked spectral fluctuations decreased with more perturbation experience for the right-side perturbations and varied with perturbation timing. Older adults showed significant electrocortical activity with a wider spread of electrocortical sources in the motor and posterior parietal cortices. Older adults demonstrated more co-contracted muscle pairs than young adults, and co-contraction did not decrease with more perturbation experience.

The results show that brief perturbation during recumbent stepping does not create error-based adaptation with reduced motor errors tied to more perturbation experience. However, these perturbations cause prolonged modifications in the motor patterns even after the perturbations are removed. Modulating the perturbation timing can tune both cortical activities at specific brain areas and modify muscular co-contraction behavior in older adults.
Dedicated to my love, Maryam.
ACKNOWLEDGMENTS

I have been very fortunate to be surrounded by friends and mentors that were kinder, smarter, more caring, and better than me throughout my life. I am forever in their debt.

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5.1 Agonist muscles to drive the stepper for the left- and right-side tasks. L = left. R = right.
CHAPTER 1: INTRODUCTION

Aims and hypotheses

Balance and walking issues are major problems with aging; >3 million older adults visit ER each year because of fall-related injuries, and >300,000 fall-related hip-fractures cost the US >$50 billion. This burden would only dramatically increase as the U.S.’s aging population will double from 57 million in 2010 to 109 million in 2050 [1]. Seated locomotor exercise, such as cycling and recumbent stepping, share similar underlying neural correlates as walking [2] but does not require maintaining balance. So, perturbed seated locomotor tasks might help the elderly be more prepared for perturbations during walking and reduce their fall risks. Recent studies have demonstrated significant improvement in walking and balance performance after unperturbed seated exercise for stroke and Parkinson’s disease patients [3, 4]. However, the motor responses and neural correlations of perturbations in seated locomotor tasks have not been studied adequately.

To determine the potential of the perturbed seated locomotor tasks with walking rehabilitation, we needed to quantify and compare biomechanical, muscular, and cortical responses during the perturbed seated task to the brain and body response to perturbation during walking. Importantly, previous studies have suggested error-based adaptation during perturbed walking. Subjects adapt their movement to reduce errors as they gain more experience with the perturbations and the adaptation would wash out soon after removing the perturbations [5]. Also, locomotor perturbations tend to increase the muscular activity and co-contraction, which would decrease with adapting to the perturbations [6, 7]. On the cortical level, the anterior cingulate cortex located at the mid-prefrontal area of the brain monitors errors during locomotor perturbations [8] and monitors walking during double support and toe-off [9, 10]. A recent study on the corticomuscular connectivity of perturbed balance-bean walking revealed a strong connection between the anterior cingulate cortex and the
motor cortex’s supplementary motor areas is present shortly after the perturbations [11].

The overarching aim of this research was to quantify the motor, muscular and cortical responses of young and older adults to locomotor perturbations during recumbent stepping, a seated arms and leg locomotor task. We aimed to contrast young and older adults’ responses to understand the possible underlying correlations of aging on human locomotion during perturbed seated locomotion. We also aimed to quantify the possible differential responses to varied perturbation timings. We had a set of hypotheses for this research: 1) Subjects would initially increase their motor errors in response to perturbations. The errors would decrease as they adapt to the perturbations. 2) Brain dynamics, especially at the anterior cingulate cortex, would reflect and monitor the presence of the perturbations. 3) Brain activity would decrease as subjects gain more experience with the perturbations. 4) Both young and older adults would increase co-contraction of the agonist and antagonist muscles in response to perturbations. 5) Young adults would benefit from more cortical activity during the perturbations, while older adults would benefit from the perturbation with greater co-contraction than young adults. 6) Young adults would have stronger corticomuscular connectivity than older adults in response to perturbations.

Research overview

We followed the research aim in two main directions: 1) We determined the accuracy and reliability of the electrocortical source estimation with the current technological advancements. Specifically, we investigated the effects of precise digitization of 3D localizing the electroencephalography (EEG) electrodes on source estimation. 2) We tested young and older adults’ cortical, muscular and biomechanical performance during perturbed recumbent stepping. Overall, 17 young adults and 11 older adults completed four perturbed stepping tasks. Each task included six minutes of perturbations during each stride, padded by two minutes of unperturbed stepping before and after
the perturbations. We perturbed the stepping movement with 200ms of increased resistance at the extension-onset or mid-extension of the left or right leg. We only used one perturbation timing in each task.

Chapters 2 and 3 are dedicated to quantifying EEG source estimation accuracy based on the reliability of the electrode locations. The second chapter deals with the variability of the different digitizing methods and how this variability can cause uncertainties in source estimation. The third chapter raises awareness about the importance of recording fiducial locations, i.e., the anatomical landmarks used to correlate the brain’s electrode locations. We emphasized that failing to record the fiducial locations accurately can result in twice as large errors in source estimation as the original error.

We discussed the biomechanics and electrocortical correlates of perturbed recumbent stepping in young and older adults in the next three chapters. We quantified the motor and electrocortical responses to perturbations in young adults in chapter 4. We discussed that the brief perturbations during recumbent stepping would not create error-based adaptation, resulting in sustained motor modifications after removing the perturbations. We could also show that the perturbations elicit theta-band (3-8Hz) synchronization (i.e., phase-locked neuronal firing) in the anterior cingulate cortex and the supplementary motor areas and that the extension-onset perturbations resulted in stronger anterior cingulate theta-band synchronization than the mid-extension perturbations.

We compared the motor error and the muscular co-contraction of young and older adults in chapter 5. We found that older adults also do not follow the error-based learning paradigm to respond to the perturbations. Young adults used a wider range of their muscles than older adults to drive the stepper during perturbations. However, older adults could reduce their co-contraction in select muscles for each task to overcome the perturbations.

Finally, we compared the electrocortical and corticomuscular correlates of perturbations between
young and older adults in Chapter 6. Unlike young adults, older adults did not present group-level anterior cingulate activity during perturbed stepping. Also, older adults’ spectral fluctuations were less pronounced in the motor cortex than young adults. Corticomuscular connectivity analysis revealed that young adults’ anterior cingulate activity directly modulates specific muscles with different perturbations. We also f was strong connectivity from the left posterior parietal cortex and older adults’ upper and lower-limb muscles around the perturbation timing.

References


CHAPTER 2: EEG ELECTRODE DIGITIZATION RELIABILITY AND SOURCE ESTIMATION UNCERTAINTY

Introduction

Estimating active cortical sources using electroencephalography (EEG) is becoming widely adopted in multiple research areas as a non-invasive and mobile functional brain imaging modality [1–4]. EEG is the recording of the electrical activity on the scalp and is appealing for studying cortical dynamics during movements and decision making due to the high temporal (i.e. millisecond) resolution of electrical signals. One of the challenges of using EEG is that the signal recorded in an EEG electrode is a mixture of electrical activity from multiple sources, which include the cortex, muscles, heart, eye, 60 Hz noise from power lines, and motion artifacts from cable sway and head movements [5, 6]. To meaningfully correlate EEG analyses with brain function, the unwanted source content such as muscle activity, eye blinks, and motion artifacts need to be attenuated or separated from the cortical signal content. A multitude of tools such as independent component analysis, artifact rejection algorithms, and phantom heads have been developed to address the need to separate the source signals to extract the underlying cortical signal [7–11]. Using high-density EEG and improving EEG post-processing techniques have also improved spatial resolution of source estimation to $\sim$ 1 cm in experimental studies [12–17].

Source estimation requires knowing the EEG signals and the locations of the EEG electrodes to estimate the locations of the cortical sources that produced the EEG signals measured on the scalp. An intuitive assumption of source estimation is that precise placement of the EEG electrodes on the scalp is essential for accurate estimation of source locations [18].

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shifts of 0.5 cm to 1.2 cm in estimated source locations as a result of 0.5 cm (or 5°) error in the electrode digitization [19–23]. For EEG studies conducted inside a magnetic resonance imaging (MRI) device, the electrode locations with respect to the cortex can be captured and processed with less than 0.3 cm position error, which results in near perfect alignment of identified brain areas [15, 24]. However, for studies that do not involve MRI, the electrode locations should be “digitized”, i.e. recorded digitally via a three-dimensional (3D) position recording method [25]. These digitized locations can then be coupled with either a subject-specific or an averaged template of the brain structure obtained from MRI or other imaging techniques to perform EEG source estimation.

Just one decade ago in the mid-2000’s, the main digitizing technologies available were based on ultrasound and electromagnetism, which were expensive, time consuming, and needed trained operators [25, 26]. An ultrasound digitizing system uses differences in ultrasound-wave travel times from emitters on the person’s face and a digitizing wand to an array of receivers to estimate the 3D location of the tip of the digitizing wand with respect to the face emitters. An electromagnetic system tracks the locations of receivers placed on the person’s head and on a wand in an emitted electromagnetic field to estimate the position of the tip of the wand with respect to the head receivers. The environment must be clear of magnetic objects when using an electromagnetic digitizing system, otherwise the electrode locations will be warped [27, 28].

Recent efforts have focused on developing technologies to make digitization more accessible and convenient, mainly by incorporating image-based technologies [29, 30]. For example, using photogrammetry and motion capture methods for digitization can provide accurate electrode locations in a short period of time [31, 32]. Photogrammetry involves using cameras to take a series of color images at different view angles. These images can then be analyzed to identify the locations of specific points in the 3D space [31, 33]. Motion capture typically uses multiple infrared cameras around the capture volume to take simultaneous images to identify the locations of reflective or
emitting markers. If markers are placed directly on the EEG electrodes, a motion capture system could conveniently record the position of all of the electrodes at once [27, 32]. Motion capture could also be used to record the position of the tip of a probe, a rigid body with multiple markers, to digitize 3D locations of the electrodes with respect to the reflective face markers. Several recent commercial digitizing systems use simple motion capture approaches to digitize EEG electrode locations with or without a probe [28, 34–36].

Another option for digitizing EEG electrodes that has also gained much interest recently are 3D scanners. A common approach for 3D scanning is detecting the infrared or visible reflections of projected light patterns with a camera to estimate the shape of an object [37]. The 3D scanned shapes can then be plotted in a software program such as MATLAB, and the locations of specific points on the 3D scanned shape can be determined. Recently, common EEG analysis toolboxes such as EEGLAB [38] and FieldTrip [39] support using 3D scanners to digitize the electrode locations. Studies suggest that 3D scanners can improve digitization accuracy and significantly reduce digitization time [40]. Using other camera-based systems such as time-of-flight scanners and virtual reality headsets were also reported to provide comparable digitization reliabilities as the ultrasound or electromagnetic digitizing methods, while reducing the time spent for digitizing the EEG electrodes [28, 41, 42].

The purposes of this study were 1) to compare the reliability and validity of five digitizing methods and 2) to quantify the relationship between digitization reliability and source estimation uncertainty. We determined source estimation uncertainty using spatial metrics and Brodmann areas. We hypothesized that digitizing methods with less reliability would increase uncertainty in the estimates of the electrocortical source locations. For our analyses, we assumed that all other contributors to source estimation uncertainty such as variability of head-meshes and assumptions of electrical conductivity values were constant.
Methods

We fitted a mannequin head with a 128-channel EEG cap (ActiveTwo EEG system, BioSemi B.V., Amsterdam, the Netherlands, Figure 2.1A) and used this mannequin head setup to record multiple digitizations of the locations of the EEG electrodes and fiducials, i.e. right preauricular, left preauricular, and nasion [43]. To prevent the cap from moving from digitization to digitization, we taped the cap to the mannequin head (Figure 2.1A). To help ensure that the fiducials were digitized at the same locations for every digitizing method, we marked the fiducials with small 4-mm markers on the mannequin head and with small o-rings on the cap (Figure 2.1A).

Digitizing methods

We compared five methods for digitization: ultrasound, structured-light 3D scanning, infrared 3D scanning, motion capture with a digitizing probe, and motion capture with reflective markers. We calibrated each digitizing device only once and completed collecting data for each digitizing in a single session. We also kept the position of the mannequin head, mannequin head orientation, start and endpoint of digitizing, lighting, and temperature constant to avoid introducing additional sources of error to our data collection and analysis.

For each method, four different members of the laboratory digitized the mannequin head five times (one person performed the digitization twice). All of the operators had prior experience in digitizing and were asked to follow each method’s specific guidelines. We imported the digitization data to MATLAB (version 9.4, R2018a, Mathworks, Natick, MA) and performed all analyses in MATLAB.
Figure 2.1: The mannequin head used for digitization and the five digitizing methods tested. **A:** The mannequin head fit with the 128-electrode EEG cap used for all of the digitizing recordings. The right and left preauriculars were marked by o-rings and nasion was marked with a reflective marker. The color-coded map of the cap shows the different electrode strips and the order of digitization from A to D. **B:** The ultrasound digitizing system and an operator placing the tip of the wand in the electrode well on the cap. Two of the total five ultrasound emitters on the face and wand, as well as the data acquisition (DAQ) box and the receiver module are also indicated. **C:** The structured-light 3D scanner and an operator manually marking the locations of individual electrodes of the scanned model in MATLAB. **D:** The infrared 3D scanner and an operator manually marking the locations of individual electrodes of the colored 3D scan in MATLAB. **E:** The motion capture digitizing probe with a close-up view of the o-rings placed 7 millimeters away from the tip. The probe has a similar role to the wand in the ultrasound system. **F:** The EEG cap with 35 3D-printed EEG electrode shaped reflective markers, 3 face markers, and 3 fiducial markers used for the motion capture digitization. We placed reflective markers on top of the preauricular o-rings to be able to capture fiducial locations. The electrode map depicts the approximate locations of the digitized electrodes and grounds.
Ultrasound

We used a Zebris positioning system with ElGuide software version 1.6 (Zebris Medical GmBH, Tübingen, Germany, Figure 2.1B) to digitize the electrodes with an ultrasound method. Following the Zebris manual, we placed 3 ultrasound emitters on the face of the mannequin head, placed the receiver module in front of the mannequin head, and used the digitizing wand to record the electrode locations. We calibrated the system using the ElGuide calibration procedure. We marked the fiducials repeatedly until we obtained fiducials with a digitized 3D location of nasion that was < 2mm with respect to the midline and with preauriculars that had a difference of < 5mm in the anterior/posterior and top/bottom directions. Operators followed the interactive ElGuide template to digitize each electrode location. This process involved fully placing the wand tip into the electrode wells and ensuring that the receivers were able to see all emitters at the time of recording electrode locations, so that the estimated position of the wand tip was stable.

Structured-light 3D scan

We used an Einscan Pro+ (Shining 3D Tech. Co. Ltd., Hangzhou, China, Figure 2.1C) to digitize the electrodes with a structured-light 3D scanner. This scanner estimates the shape of an object from reflections of the projected visible lights. We calibrated the Einscan Pro+ one time with the Einscan’s calibration board and followed the software’s step-by-step instructions. We used the scanner’s hand-held rapid mode with high details and allowed the scanner to track both texture and markers during the scanning process. Each operator scanned the mannequin head until the scan included the cap, fiducials, and the face. We then applied the watertight model option to the scan and exported the model as a PLY file to continue the digitization process in MATLAB.

After acquiring the 3D scan, the 3D head model needed to be imported into a software program,
where the operator manually marked the EEG electrode locations on the 3D scanned head model. We followed the FieldTrip toolbox documentation for digitization using 3D scanners [44] and created a MATLAB script file for importing and digitizing 3D models of the mannequin head. The operator first marked the fiducials on the mannequin head model in MATLAB to build up the head coordinate system. Then, the operator marked the locations of the electrodes on the screen in each section of the cap in alphanumerical order (A, B, C, D, and the fiducials, total: 131 locations, Figure 2.1A). The operators referred to a physical EEG cap for guidance to help mark the locations in the expected order because these scans were not in color and the letter labels of the electrodes were not visible on the 3D model.

Infrared 3D scan

We used the Structure sensor (model ST01, Occipital Inc., San Francisco, CA) integrated with an Apple® iPad (10-inch Pro) to digitize the electrodes with an infrared dot-projection 3D scanner (Figure 2.1D). This scanner shares similar working principles as a structured-light scanner but uses infrared light projection to estimate the shape of objects. We calibrated the sensor in daylight and office light according to the manual. We scanned the head using the high color and mesh resolutions. When the mannequin head was completely in the sensor’s field of view, the operator started scanning. The Structure sensor interface gives the operator visual feedback to help the operator obtain a complete high-quality scan. We visually inspected that the scanned model matched the mannequin and then exported the model to the MATLAB environment. We used the FieldTrip toolbox to import and digitize the 3D mannequin head scans following the same procedure described for the structured-light 3D scan digitization.
**Motion capture probe**

We used a digitizing probe and a 22-camera motion capture system (OptiTrack, Corvallis, OR) to digitize the electrodes. The probe is a solid rigid body with four fixed reflective markers (Figure 2.1E). We placed three reflective makers on the face of the mannequin to account for possible movements of the head during data collection. Each operator digitized the fiducials and each section of the cap (A, B, C, D, Figure 2.1A) in separate takes. We placed double o-rings seven millimeters away from the probe tip to ensure consistent placement of the tip inside the electrode wells (Figure 2.1E). The tracking error of the motion capture system was less than 0.4 millimeters.

**Motion capture**

We used the motion capture system to record the locations of 35 3D printed reflective markers that resembled a 4-millimeter reflective marker on top of a BioSemi active pin electrode (Figure 2.1F). We did not use actual BioSemi electrodes, which have wires that could prevent the cameras from seeing the markers. We placed 27 EEG electrode shaped markers to approximate the international 10-20 EEG cap layout and placed an additional eight EEG electrode shaped markers randomly on the cap to add asymmetry to improve tracking of the markers. We recorded 2-second takes of the positions of the 35 markers, three markers on the fiducials, and three face markers. Before transforming the locations to the head coordinate system, we identified and canceled movements of the head during data collection using the three face markers.

**Transformation to head coordinates**

We developed a dedicated pipeline to convert the digitized electrode locations for each digitizing method to a format that could be imported to the common toolboxes for EEG analyses. Because
EEGLAB and FieldTrip can easily read Zebris ElGuide’s output file (an SFP file), we created SFP files for all digitizations.

The head coordinate system in ElGuide defines the X-axis as the vector connecting the left preauricular to the right preauricular and the origin as the projection of the naison to the X-axis. Therefore, the Y-axis is the vector from the origin to the naison, and the Z-axis is the cross product of the X and Y unit vectors, which starts from the origin.

**Digitization reliability and validity**

Variations in the digitized electrode locations could originate from random errors and systematic bias. The effects of random errors can be quantified as variability. Reliability is also inversely related to variability. Systematic bias can be quantified as the difference between measured locations and the ground truth locations. Validity is also inversely related to systematic bias.

**Digitization reliability**

To assess the effects of random errors, we quantified digitization variability. We averaged the five digitized locations for each electrode to find the centroid. We then calculated the average Euclidean distances of the five digitized points to the centroid for each electrode and averaged those distances for all of the electrodes to quantify within-method variability. We identified and excluded outliers, single measurements that were beyond five standard deviations of the average variability for a digitizing method (1 out of 655 measurements for ultrasound, 4 out of 655 measurements for motion capture probe and 2 out of 190 measurements for motion capture). If there were outliers, we recalculated the average digitization reliability with the updated dataset. Throughout the paper, we use “variability” to refer to “within-method variability.” Because reliability is inversely related to
the variability, the most reliable method has the least variability.

**Digitization validity**

To quantify the systematic bias of a digitizing method, we calculated the average Euclidean distance between the centroid for a digitizing method and the ground-truth centroid for the same electrode. We used the electrode centroids from the most reliable digitizing method as the ground-truth [45]. Then, we averaged the Euclidean distances for the 128 electrodes to obtain the magnitude of the systematic bias for each digitizing method. Because validity is inversely related to the systematic bias, the most valid method has the least systematic bias.

**Source estimation uncertainty**

To generalize the possible effects of digitization reliability, we synthesized 500 sets of electrode locations with a Gaussian distribution using the variability average and standard deviation calculated for each digitizing method in 2. We excluded the motion capture method from the source estimation uncertainty analyses because we only recorded the locations of 35 EEG electrode shaped reflective markers instead of all 128 EEG electrode locations. We used a single representative 128-channel EEG dataset from a separate study for the source estimation analyses. We applied the Adaptive Mixture Independent Component Analysis (AMICA) to decompose EEG signals into independent components (ICs) [46], which has been reported to represent dipolar activities of different brain and non-brain sources [7].

We used EEGLAB’s DIPFIT toolbox version 2.3 to estimate a dipole equivalent for each IC and applied DIPFIT 500 times for each digitizing method. Each DIPFIT iteration used one of the 500 sets of synthesized electrode locations, the Montreal Neurological Institute (MNI) head model [47],
and the ICs from the AMICA. The MNI head model is an averaged structural head model from 305 participants and provides 1mm × 1mm × 1mm resolution. To convert the mannequin head to be compatible with the MNI model, we warped the electrode locations to the MNI model using only the fiducials to preserve individual characteristics of the mannequin head. We used the dipoles produced with the electrode location centroids from the digitizing method with the highest reliability and identified the dipoles that described > 85% of the IC signal variance. We also excluded any dipole that was estimated to be outside of the brain volume for any of the DIPFIT results (500/method × five methods = 2500 DIPFIT results). In the end, 23 ICs remained.

**Spatial uncertainty**

We fitted an enclosing ellipsoid with the minimum volume to each IC’s cluster of 500 dipoles [48] and quantified spatial uncertainty in terms of the volume and width of the ellipsoid. A larger ellipsoid volume indicated that a single dipole could reside within a larger volume, and thus, had greater volumetric uncertainty. A larger ellipsoid width indicated that a single dipole could have a larger shift in location. We calculated the ellipsoid’s width as the maximum distance that the IC’s dipoles could have from one another. We averaged the volumes and widths of all 23 ICs to quantify the spatial uncertainty for each digitizing method.

**Brodmann area accuracy**

To identify Brodmann areas, we used a modified version of the eeg_tal_lookup function from EEGLAB’s Measure Projection Toolbox (MPT). This function looks for the anatomic structures and Brodmann areas in a 10-mm vicinity of each dipole and assigns the dipole to the Brodmann area with the highest posterior probability [49, 50]. We identified the “ground-truth” Brodmann areas from the dipoles estimated using the centroid electrode locations of the most reliable digitizing
method. Then, we calculated Brodmann area accuracy as the percentage of the other 500 Brodmann area assignments that matched the “ground-truth” Brodmann area.

We also analyzed Brodmann area accuracy using a template of electrode locations based on the MNI head model [51]. Because the BioSemi 128-electrode cap is not based on the 10-10 electrode map, instead of using the 10-10 electrode locations, we warped the BioSemi electrode locations to reside on the outer surface of the MNI head model. We then compared the Brodmann area identified from the template to the “ground-truth” Brodmann area. Since there is only one template for the Biosemi 128-electrode location on the MNI head model and the locations are fixed, we could not calculate a percentage of assignments; thus, the template’s Brodmann area for each IC was either a hit or miss. However, we did calculate and compare the distance between the template’s dipole to the “ground truth” dipole. We also compared the distance between each digitizing method’s dipoles to the “ground truth” dipole. These distances indicated whether the dipoles estimated using each digitization method were near the “ground truth” dipole.

**Statistical analysis**

We used a one-way repeated measures analysis of variance (rANOVA) to compare the reliability and validity of the digitizing methods, the spatial uncertainty of the estimated dipoles, and the Brodmann area accuracy. For significant rANOVA’s, we performed Tukey-Kramer’s post-hoc analysis to determine which comparisons were significant. We also performed a one-sided Student t-test to identify if the Brodmann area accuracy of each digitizing method was different from the template. The level of significance for all statistics was $\alpha = 0.05$. For rANOVA, we reported degrees of freedom (DF), Fisher’s F-test result and the probability value (p-value). We used p-values to report post-hoc and Student t-test results.

Additionally, we fit a polynomial, using a step-wise linear model (MATLAB `stepwiselm` function),
to describe spatial uncertainty as a function of digitization variability. We forced the y-intercept of
the first-order polynomials and the y-intercept and y’-intercept of the higher-order polynomials to be
zero. We set the y-intercepts to be zero for two reasons: 1) when we used the exact same electrode
locations and performed DIPFIT 100 times, the maximum distance between source locations was
on the order of $10^{-4}$ cm, and 2) the fit should not model the uncertainty values $< 0$ for positive
digitization variability values. The step-wise linear model started with a zero order model and only
added a higher-order polynomial term when necessary. The criterion for adding a higher-order
polynomial term to the model was a statistically significant decrease of the sum of the squared error
between the data points and the predicted values.

Results

The variability results for the five digitizing methods were visibly different, and electrodes located at
the back of the head tended to have greater variability (Figure 2.2). The variability for the ultrasound
method was generally largest compared to the other methods and could be as large as $\sim 1.5$ cm
for electrodes at the back of the head. The variability for all electrodes digitized with the motion
capture method was small, being no greater than 0.001 cm.

There was a range of reliabilities among the digitizing methods (Figure 2.3A). The motion capture
digitizing method had the smallest variability of $0.001 \pm 0.0003$ cm (mean $\pm$ standard deviation)
and hence, the greatest digitization reliability. The motion capture probe was the next most reliable
method with an average variability of $0.147 \pm 0.03$ cm, followed by the infrared 3D scan ($0.24
\pm 0.05$ cm), the structured-light scan ($0.50 \pm 0.09$ cm), and the ultrasound digitization ($0.86 \pm
0.3$ cm). The variability for the digitizing methods were significantly different (rANOVA DF=4,
$F=1121$, $p<0.001$), and the variability for each digitizing method was significantly different from
all other digitizing methods (post-hoc Tukey-Kramer, $p$’s $<0.001$).
Figure 2.2: Visualization of the digitization reliability. Colored and scaled dots show the electrode location within-method variability for all 128 electrodes for the five digitizing methods. Ultrasound had the greatest variability and was the least reliable. The electrodes at the back of the head also tended to have the greatest variability. The motion capture method had the least variability and was the most reliable. The color bar and scale for the radii of the dots illustrate the magnitude of variability.

The systematic biases, thus validities, of the digitizing methods were significantly different (rANOVA DF=2, F=143.1, p<0.01, Figure 2.3B). The digitization validity of the structured-light 3D scan was the worst of the digitizing methods with a systematic bias of 0.63 ± 0.18 cm that was significantly larger than the other digitizing methods (post-hoc Tukey-Kramer, p’s <0.001). The digitization validity of the ultrasound and the infrared 3D scans were similar, with systematic biases of 0.43 ± 0.18 cm and 0.41 ± 0.13 cm, respectively.
Figure 2.3: A: Reliabilities, quantified as the average variability, were significantly different for the five digitizing methods. The reliability of each digitizing method was significantly different from all other methods (* = Tukey-Kramer p’s < 0.001 for all pair-wise comparisons). B: Validity, quantified as the average systematic bias showed that the structured-light 3D scan had the largest systematic bias compared to ultrasound and the infrared 3D scan. The motion capture probe method was assumed to be the ground truth and thus has no systematic bias and is not shown. * = Tukey-Kramer p’s < 0.001. Error bars are the standard deviation. infrared = infrared 3D scan. str.-light = structured-light 3D scan. m.+probe = motion capture probe. mocap = motion capture.

Within a given digitizing method, dipoles generally showed similar spatial uncertainty while different digitizing methods generally showed differences in spatial uncertainty (Figure 2.4). Ellipsoid sizes for the motion capture probe, infrared 3D scan, structured-light 3D scan, and ultrasound digitization increased in order from the smallest to the largest, respectively. The enclosing ellipsoids of adjacent
ICs also overlapped when the ellipsoid size was large, on the order of 1 cm$^3$, such as for the ultrasound method.

Figure 2.4: An example depiction of the synthesized electrode locations with a Gaussian distribution using the same averaged variability and standard deviation as the structured-light 3D scans, and the enclosing ellipsoids of the 500 dipoles for each independent component (IC) and digitizing method. Black dots = centroids of the electrode locations. Light gray dots = first 150 out of 500 synthesized electrode locations. Each color represents a different IC (23 ICs total). A close-up view of the ellipsoid fit for an Anterior Cingulate IC based on the reliability of the ultrasound digitizing method.

Ellipsoid volumes increased significantly with increasing digitization variability among the digitizing methods and had a cubic relationship ($r^2 = 1.00$, Figure 2.5A). The motion capture probe and infrared 3D scan had the smallest uncertainty volumes (mean ± standard error) 0.007 ± 0.0007 cm$^3$ & 0.029 ± 0.0027 cm$^3$, respectively, whereas ultrasound had the largest uncertainty volume (1.37 ± 0.13 cm$^3$). Structured-light 3D scan had an average uncertainty volume of 0.21 ± 0.014 cm$^3$. The volumes of the enclosing ellipsoids showed a significant between-group difference (rANOVA,
DF=3, F=114.4, p<0.001), and all uncertainty volume combinations of paired digitizing methods were significantly different (Tukey-Kramer post-hoc, p’s<0.001).

Ellipsoid widths also increased significantly with increasing digitization variability among the digitizing methods but had a linear relationship where the ellipsoid width was twice the size of the digitization variability ($r^2 = 1.00$, Figure 2.5B). The average ellipsoid width was the smallest for the motion capture probe, (mean ± standard error) 0.34 ± 0.018 cm. The average ellipsoid widths for the two 3D scans were 0.53 ± 0.028 cm for the infrared 3D scan and 1.09 ± 0.051 cm for the structured-light 3D scan. The largest average ellipsoid width was for the ultrasound digitization, 1.90 ± 0.081 cm. The rANOVA for the widths of the enclosing ellipsoids showed a significant between-group difference (DF=3, F=434.8, p<0.001) and all combinations of paired digitizing methods had significantly different uncertainty widths (Tukey-Kramer post-hoc p’s<0.001).

The Brodmann area accuracy among the digitizing methods could be extremely consistent within some ICs and could also be drastically different for other ICs (Figure 2.6 and in Supplement Figure S1). In general, the digitizing method with the highest reliability also had the highest Brodmann area accuracy within a given IC. For some ICs, all digitizing methods had > 98% Brodmann area accuracy. For other ICs, the Brodmann area accuracy decreased as reliability decreased. The most drastic example for this dataset was BA18 in Figure 2.6, where the Brodmann area accuracy was 86% with the motion capture probe method but dropped to 26% with the ultrasound method.

The Brodmann area accuracy for the digitizing methods and the template were significantly different (Figure 2.7). The motion capture probe had the highest Brodmann area accuracy, 93% ± 16 (mean ± standard deviation). The remaining digitizing methods in order of decreasing Brodmann area accuracy were the infrared 3D scan (91% ± 19%), the structured light 3D scan (87% ± 23%), and the ultrasound digitization (79% ± 25%). The rANOVA for the Brodmann area accuracy showed a significant between-group difference (DF=4, F=306.4, p<0.001). Post-hoc Tukey-Kramer analysis
showed significant pair-wise difference between all groups except the motion capture probe and infrared 3D scan. Using the MNI electrode template decreased the Brodmann area accuracy to 53% and was significantly different compared to any of the digitizing methods (p’s < 0.001). The average distance of the dipoles of each digitizing method to the “ground-truth” dipole was less than 0.4 cm while the average distance of the template dipoles to the “ground-truth” dipole was ∼ 1.4 cm.

Figure 2.5: The relationships between digitization variability and dipole spatial uncertainty. A: Digitization variability and ellipsoid volume had a cubic relationship with an $r^2$ of 1.00. B: Digitization variability and ellipsoid width had a linear relationship with an $r^2$ of 1.00. Error bars are the standard error. * = Tukey-Kramer p’s <0.001 for all pair-wise comparisons. m.+probe = motion capture probe. infrared = infrared 3D scan. str.-light = structured-light 3D scan.
Discussion

We found that there was a range of reliability and validity values among the digitizing methods. We also observed that less reliable digitizing methods translated to greater uncertainty in source estimation and poorer Brodmann area accuracy, assuming all other contributors to source estimation uncertainty were constant. Of the five digitizing methods (ultrasound, structured-light 3D scan, infrared 3D scan, motion capture probe, and motion capture), the most reliable digitizing method was the motion capture while ultrasound was the least reliable. The structured-light digitizing method had the greatest systematic bias and was thus the least valid method. We had hypothesized that less reliable digitizing methods would lead to greater source estimation uncertainty. In support of our hypothesis, digitizing methods with decreased reliability resulted in increased spatial uncertainty of the dipole locations and decreased Brodmann area accuracy. Surprisingly, any digitizing method led to an average Brodmann area accuracy of $>80\%$. Using a template of electrode locations decreased Brodmann area accuracy to $53\%$. Overall, these results indicate that electrode digitization is crucial for accurate Brodmann area identification using source estimation and that more reliable digitizing methods are beneficial if the functional resolution for interpreting source estimation is more specific than Brodmann areas.

To help summarize the advantages of the different digitization systems, we created a table comparing the digitization reliability, dipole uncertainty, speed, affordability, and ease-of-use score, which are different factors that could influence which digitization a laboratory might choose to use (Table 2.1). We estimated the digitizing speed as how much time each digitization required. The fastest digitizing method that required manual electrode marking was the motion capture probe method, which took 5 minutes to mark each electrode and 5 minutes to calibrate the system. The least expensive system was the infrared 3D scanner, which is likely to become even less expensive as cameras on smartphones become more advanced and could soon be used to obtain an accurate 3D
Figure 2.6: Brodmann area (BA) accuracy for a subset of ICs. The dipole depicts the “ground truth” dipole produced from the most reliable digitizing method, the motion capture probe method. The pie charts show the distribution of the Brodmann area assignments compared to the “ground truth” Brodmann area (shown in bold). ICs in the left column had consistent Brodmann area assignments regardless of digitizing method while the ICs in the right column had more varied Brodmann area assignments for the different digitizing methods. In general, less reliable digitizing methods led to less consistent Brodmann area assignments.

scan for digitizing EEG electrodes. We also surveyed the operators to score each digitization on a scale of 1-5, with 1 being easy to use. While performing the actual 3D scan was perceived as being easy, marking the electrodes in MATLAB was not an easy task. The operators indicated that the motion capture was the easiest and that ultrasound was the most difficult method to use. To create a final ranking, we averaged the rankings for each factor (digitization reliability, dipole uncertainty, speed, affordability, and ease-of-use) to obtain a method score. Based on the method score, the
best digitizing method was the motion capture. The next best method was tied between the motion capture probe and infrared 3D scan. The fourth best digitizing method was the structured-light 3D scan, and the worst digitizing method was the ultrasound method, which ranked poorly for all factors.

Our results suggest that the motion capture method currently provides the most reliable electrode digitization. The average variability of the motion capture digitization was less than the mean calibration error reported by the motion capture system (0.001 versus <0.04 cm respectively). This difference might be because of the different natures of the two variabilities. The digitization variability is defined for a seated subject (or mannequin) and multiple sub-second snapshots of the static electrodes placed on the cap. However, the mean calibration error is defined for a set of moving markers in a much larger volume across several minutes of a calibration period. Using the same position for mannequin placement and lack of head movement may have also contributed to the small digitization variability using motion capture. In a previous study, Reis and Lochmann developed an active-electrode motion capture approach for an EEG system with 30 electrodes and reported small deviation of the digitized locations from the ground truth locations [32]. In addition to having sub-millimeter variability, the motion capture method only required 1-2 seconds to digitize, assuming that the markers were already placed on the EEG electrodes. However, tracking 64+ markers on an EEG cap may be challenging for most motion capture systems. Determining the maximum number of EEG electrodes that could be digitized using a motion capture approach could be beneficial and pursued in future work. Laboratories that already have a motion capture system and do not need to digitize more than 64 EEG electrodes could conveniently use the motion capture method, which would provide a cost-effective, fast, and easy digitizing process. For laboratories that need to digitize 64+ electrodes and have a motion capture system already, the motion capture probe digitizing method would be the recommended option.

Our results support recent efforts to use 3D scanners as a reliable and cost-effective method to
digitize EEG electrodes [40, 52, 53]. Both the structured-light and infrared 3D scanning methods were more reliable than digitizing with the ultrasound method. Furthermore, our reliability results for the two 3D scanners align well with a recent study that showed that an infrared 3D scan could automatically digitize electrode locations on three different EEG caps and achieve good reliability after additional post-processing [40]. Of the two 3D scanners we tested, the less expensive infrared 3D scanner was more reliable, had higher validity, and resulted in less dipole uncertainty, compared to the structured-light 3D scanner. Even though the structured-light 3D scanner provides more details from the mannequin head and cap, those details did not seem to be important for improving digitization reliability or validity. Additionally, the highly detailed structured-light 3D scans created large files and resulted in sluggish refresh rates that made using FieldTrip toolbox to rotate and manipulate the 3D scans difficult. The infrared 3D scan, unlike the structured-light 3D scan, was in color, which was helpful for the operators to identify the EEG electrodes more easily on the computer screen. In the future, artificial intelligence approaches may be able to fully automate the digitizing process and use the additional topographic details from high resolution 3D scans. A continuous image-based digitizing method such as using a regular video recorded using a typical smartphone could also potentially be developed to digitize EEG electrode locations.

Compared to simulation studies, our experimental results demonstrated that source estimation uncertainty increased steeply with increasing EEG electrode variability. We showed that a digitizing method with an average variability of 1 cm could lead to a shift of a single dipole by more than 2 cm, which is $>20\%$ of the head radius. There is just one simulation study that we know of that also showed a 2-fold increase in source uncertainty for every unit of digitization variability [23]. In that study, digitization variabilities were created using systematic rotations applied to every electrode location. The majority of the simulation studies however, suggest that source uncertainty could only be as large as the digitization variability [20–22, 54]. In one of the mathematical studies, the theoretical lower bound of source estimation uncertainty was 0.1 cm for 0.5 cm shifts in EEG
electrode location [22], which is 10x smaller than our experimental results. While simulation studies can be insightful, results should also be cross-validated with a conventional source estimation method (e.g. DIPFIT, LORETA or minimum norm) to determine whether simulation results are indicative of real-world source estimation uncertainty.

Because researchers often use Brodmann areas to describe the function of a source, we translated our results to be in terms of Brodmann area accuracy, which led to a few surprising revelations. The main revelation was that despite the range of digitization reliabilities, any of the digitizing methods we tested produced an average Brodmann area accuracy $> 80\%$. As long as sources are only discussed according to Brodmann areas or larger cortical spatial regions, any current digitizing method can be used. The second revelation was that using the template electrode locations, instead of digitizing the electrodes, significantly decreased Brodmann area accuracy from $> 80\%$ to $\sim 50\%$, which may be due to a $\sim 1.5$cm shift in dipoles locations (Figure 2.7). This shift may occur because the template removes information related to individual’s head shape. The third revelation was that for several sources, the same Brodmann area was almost always identified, regardless of the digitizing method used (left column in Figure 2.6). For other sources, less reliable digitizing methods led to more potential Brodmann area assignments (right column in Figure 2.6), but those different Brodmann areas may be functionally similar. Most likely, the proximity of a source to the boundary of a Brodmann area as well as the size of the Brodmann area contribute to Brodmann area accuracy. Ultimately, the accuracy of source estimation will depend on the target volumes of cortical regions of interest.
Figure 2.7: Brodmann area accuracy plotted versus the average dipole distance from the “ground truth” dipole when using different digitizing methods and the MNI template. Because larger distances between the dipoles and the “ground truth” likely would decrease Brodmann area accuracy, we plotted the methods on the x-axis at the method’s averaged dipole distance from the “ground truth” dipole. The box-whisker plot contains the Brodmann area accuracy averages for the 23 ICs. The Brodmann area accuracy average for an IC was the average of the percentage of the 500 iterations when the Brodmann area identified matched the “ground truth” Brodmann area for that IC. For the template, 53% of the Brodmann areas assigned for the 23 ICs using the template matched the “ground truth” Brodmann area. The Brodmann area accuracy was significantly different among the digitizing methods, except between the motion capture probe and infrared 3D scan (* Tukey-Kramer p’s <0.001). The template’s Brodmann area accuracy was significantly different than all digitizing methods (# Student’s t-test p’s <0.001). m.+probe = motion capture probe. infrared = infrared 3D scan. str.-light = structured-light 3D scan.

This study does not account for all of the possible sources of errors contributing to digitizing EEG electrodes or source estimation. We placed markers on the fiducials to control for the digitization error of the fiducials, but in practice, marking the fiducials while the subject wears the cap can be challenging. Mismarking a fiducial can significantly shift every dipole location by 2 times the distance of the fiducial mismarking [55]. We also used a mannequin to control for the head movements and relative cap movements to the head. In reality, participants may move their head...
and the cap may slightly change position during digitization or data collection that would affect the location of the EEG electrodes with respect to the head. Further, we only calibrated our digitizing devices once for multiple data collections. Nevertheless, in a real laboratory setup, device calibration might be required before each instance of data collection. We, however, included digitization by multiple experienced operators to acknowledge that in a research laboratory different members might complete the digitization for different participants. Overall, our results suggest that as long as all sources of digitization error do not create variability > 1 cm, Brodmann area accuracy would be > 80%. Using the same electrical head model and source localization approach helped us to only quantify the effects of digitization variability on source estimation uncertainty. In reality, the EEG signal noise, number and distribution of EEG electrodes, electrical properties of the head model, head model shape and mesh accuracy, and solving approach are among the other potential contributors to source estimation uncertainty [23, 45, 56–59].

Limitations of this study were that we tested a subset of all digitizing methods, used a mannequin head, and not an actual human head for the digitization, and did not perform source estimation using other common algorithms. Even though we did not test many of the marketed digitizing systems, we replicated and tested the fundamental methods used by most of the marketed digitizing systems. One widely used EEG electrode digitizing method we did not test is an electromagnetic digitizing method (e.g. Polhemus Patriot or Fastrack system). Another study using similar digitization reliability analyses reported an average variability of 0.76cm for an electromagnetic digitization system [31], which is slightly better than the ultrasound digitizing method, with a variability of 0.86 ± 0.3cm. Collecting digitization data from an actual participant might have helped in having a better distribution of the sources inside the brain volume, but we decided to use a mannequin to better control for head movements, relative cap movements and other environmental factors. Here, we used the EEG data only to provide a platform to understand the relationship between the digitization variability and source uncertainty, and locations of the sources do not have any
neurological implications. Last, we did not use other different source estimation algorithms such as LORETA or beam-forming. Studies indicate that commonly used source estimation algorithms generally identify the similar source locations [57, 58, 60], which suggests that the choice of the source estimation algorithm used would probably not significantly alter our results.

Table 2.1: Rankings for each digitizing method based on factors related to performance, cost, and convenience. # is the rank of each method among all five methods and for the specified factor. The digitization reliability values and dipole uncertainty scalar width values were taken from our results. Speed was the approximate time a digitizing method required to obtain the file of electrode locations. The ease-of-use score was the average score operators provided in a survey with a score of 5 being the most difficult and 1 being the easiest method to do. The method score is the average rank of all factors for a given method and was defined as $score = \sum \# / N$. Dipole uncertainty was not available for motion capture digitization. Mocap = motion capture. Str.-Light 3D = structured-light 3D scan. * The probe price is for the OptiTrack digitizing probe. ** motion capture cost was for an eight-camera system (Optitrack Flex13, $8000) and the Optitrack Motive software ($3000).

<table>
<thead>
<tr>
<th>method</th>
<th>digitization reliability</th>
<th>dipole uncertainty</th>
<th>speed</th>
<th>affordability</th>
<th>ease-of-use</th>
<th>method score</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mean ± SD (cm)</td>
<td>mean ± SD (cm)</td>
<td>time</td>
<td>cost (USD)</td>
<td>(1 is easiest)</td>
<td>(lower is better)</td>
</tr>
<tr>
<td>ultrasound</td>
<td>0.86 ± 0.30</td>
<td>1.90 ± 0.39</td>
<td>20</td>
<td>15k</td>
<td>4.2</td>
<td>4.6</td>
</tr>
<tr>
<td>infrared 3D scan</td>
<td>0.24 ± 0.05</td>
<td>0.53 ± 0.136</td>
<td>3</td>
<td>1k</td>
<td>2.5</td>
<td>2.3</td>
</tr>
<tr>
<td>str.-light 3D scan</td>
<td>0.50 ± 0.09</td>
<td>1.09 ± 0.24</td>
<td>25</td>
<td>5k</td>
<td>3.4</td>
<td>3.3</td>
</tr>
<tr>
<td>mocap probe</td>
<td>0.15 ± 0.03</td>
<td>0.34 ± 0.08</td>
<td>10</td>
<td>1k$^*$ (probe)</td>
<td>2</td>
<td>2.2</td>
</tr>
<tr>
<td>mocap</td>
<td>0.001 ± 0.0003</td>
<td>N/A</td>
<td>5.1</td>
<td>11k$^{**}$</td>
<td>3</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Future efforts to improve source estimation, so that sources can be interpreted in terms of cortical spatial regions smaller than Brodmann areas, will involve more than just developing more reliable, convenient, and cost-effective digitizing methods to help reduce source estimation uncertainty. Even if a perfect digitizing method could be developed, there would still be uncertainty in source
estimation as result of other factors such as improper head-model meshes and inaccurate electrical conductivity values [56, 59], which were assumed to have a constant contribution to the source estimation uncertainty in our analyses. Obtaining and using as much subject specific information, such as subject-specific MRI scans in addition to digitizing EEG electrode locations, should improve source estimation. EEGLAB’s Neuroelectromagnetic Forward Head Modeling Toolbox (NFT) could be used to warp the MNI head model to the digitized electrode locations to retain the individual’s head shape but is computationally expensive [61]. Using subject-specific MRIs instead of the MNI head model is also limited to groups with access to an MRI at an affordable cost per scan.

Conclusion

In summary, there was a range of digitization reliabilities among the five digitizing methods tested (ultrasound, structured-light 3D scanning, infrared 3D scanning, motion capture with a digitizing probe, and motion capture with reflective markers), and less reliable digitization resulted in greater spatial uncertainty in source estimation and poorer Brodmann area accuracy. We found that the motion capture digitizing method was the most reliable while the ultrasound method was the least reliable. Interestingly, Brodmann area accuracy for a source only dropped from ∼ 90% to ∼ 80%, when using the most and least reliable digitizing methods, respectively. If source locations will be discussed in terms of Brodmann areas, any of the digitizing methods tested could provide accurate Brodmann area identification. Using a template of EEG electrode locations, however, decreased the Brodmann area accuracy to ∼ 50%, suggesting that digitizing EEG electrode locations for source estimation results in more accurate Brodmann area identifications. Even though digitizing EEG electrodes is just one of the factors that affects source estimation, developing more reliable and accessible digitizing methods can help reduce source estimation uncertainty and may allow sources to be interpreted in terms of cortical regions more specific than Brodmann areas in the future.
References


CHAPTER 3: INFLUENCE OF MISMATCHING FIDUCIALS ON EEG SOURCE ESTIMATION

Introduction

Digitizing electrode locations is an essential step for setting up a head model to estimate cortical and subcortical sources from magneto-/electro-encephalography (M/EEG) signals [1]. The digitizing process involves recording three-dimensional positions of the M/EEG electrodes in a global coordinate system and transforming the locations from the global coordinates to the head coordinate system. This transformation requires that the two coordinate systems share at least three anatomical landmarks (i.e. fiducials). The fiducials are typically the left preauricular, right preauricular, and nasion [1, 2].

After digitizing the 3D locations of the fiducials and electrodes, these locations are warped to a head model or vice versa [3]. For studies that involve concurrent tomographic imaging such as magnetic resonance imaging (MRI), digitization, transformation to the head coordinate system, and warping to the head model can be made simultaneously [4], but for other studies, digitized locations must be manually coregistered to construct the head model [1]. Therefore, digitizing can significantly affect the ability to achieve a realistic head model and estimations of source locations [3].

To our knowledge, only a couple of studies examined the effects of digitizing errors on source estimation, and outcomes of these studies do not seem consistent. Beltarchini and colleagues [5] suggested that effects of electrode mislocations are negligible on the estimated source locations, while Dalal et al. [6] showed that quality of the output signal degrades significantly with higher

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1This chapter has been published in bioRxiv, and was presented in the 2019 IEEE/EMBS conference on Neural Engineering (NER), doi:10.1101/544288
uncertainties in the electrode digitization. We did not find any published study that examined the possible effects of mismarking fiducial locations on source estimation.

The purpose of this study was to analyze changes in the estimated source location as a result of shifting the fiducial locations to simulate mismarking of the fiducials. We hypothesized that changes in the locations of the fiducials would have a significant effect on the dipole fitting process and source localization.

Methods

We fitted a mannequin head with a 128-electrode BioSemi (BioSemi B.V., Amsterdam, the Netherlands) cap and digitized the locations of the electrodes and fiducials (left preauricular, LPA; right preauricular, RPA; and nasion, Nz) using an OptiTrack 22-camera motion capture system and a digitizing probe (NaturalPoint Inc., Corvallis, OR). The average (±SD) reliability of this digitization method was <1.50 ± 0.3 mm for digitizing the mannequin head five times.

To define the head coordinate system, we assumed the LPA and RPA were on the X-axis, and Nz was on the Y-axis (Figure 3.1A). The origin of the head coordinate system was located at the projection of Nz to the X-axis. The Z-axis was defined as the cross product of the X and Y unit vectors and began from the origin. We transformed the digitized electrode locations of the mannequin head to the head coordinate system and used this set of electrodes as the baseline.

To create multiple sets of electrode locations with various fiducial locations, we shifted one fiducial at a time, in 1 mm increments, up to ±20 mm in the Y and then the Z directions for the preauriculars, RPA and LPA, and in the X and then the Z directions for the nasion, Nz. This process resulted in 12 fiducial shifts per 1 mm increment (3 fiducials × four directions), totaling 240 sets of electrode locations (12 fiducial shifts per 1 mm increment × 20 fiducial shift increments).
Figure 3.1: A. The mannequin head used for digitization with a representation of the head coordinate system. Right preauricular (RPA) and nasion (Nz) are labeled in the picture. B. The baseline dipole locations of the 23 independent components (ICs) from a separate study but estimated with the mannequin head digitization rather than the individual subject digitizations using ultrasound. These ICs were used to analyze the influence of the systematic fiducial shifts on the dipole locations.

Digitization post-processing and the subsequent analyses throughout the study were performed using MATLAB version 9.4 (R2018a, Mathworks Inc., Natick, MA) and EEGLAB [7] version 14.1.2. We used just the fiducial locations to warp the locations of the electrodes and fiducials to the Montreal Neurological Institute’s template head model.
Source Estimation and Dipole Fitting

We used a single representative EEG dataset and weightings of a single Adaptive Mixture Independent Component Analysis (AMICA) from a separate study to perform dipole fitting using the multiple sets of electrode locations with the systematic shifts of the fiducial locations. This representative EEG dataset and ICA had 89 independent components, ICs. We used the DIPFIT toolbox version 2.3 to fit dipoles to the ICs. We visually inspected for dipoles with residual variances <15% that stayed inside the brain area across every fiducial shifts, which resulted in total 23 remaining ICs (Figure 3.1B). We analyzed the influence of the fiducial shifts on all twenty-three ICs but picked three to illustrate dipole location alterations in detail: 1) the anterior cingulate, 2) the primary somatosensory cortex, and 3) the premotor cortex. Previous studies have shown that these three areas are active in locomotion and error monitoring [8, 9].

Dipole Uncertainty Analysis

We defined the baseline set of dipoles as the dipoles produced using the baseline locations for the fiducials and electrodes (i.e. no shifts). Each set of electrode locations based on a shifted fiducial produced a new set of dipoles. There were 240 sets of electrode locations resulting in 240 sets of dipoles around the baseline dipole. We considered that the spread or size of the cluster of dipoles was representative of the uncertainty related to the resulting dipole locations.

We quantified dipole uncertainty as the volume of a set of tetrahedrons formed from connecting the 12 dipoles created by shifting every fiducial in 1-mm step in every direction (three fiducials × 4 direction/fiducial). We identified the outside boundary of the 12-dipole clusters and created the tetrahedrons using MATLAB’s convexHull function.

We also calculated the maximum dipole cluster width between the dipoles with equidistant fiducial
shifts (i.e. the same 12 dipoles used for creating tetrahedrons). Since the tetrahedral volumes did not have similar shapes at each fiducial shift, we formed equivalent rectangular cubes with volumes equal to the tetrahedral volumes and the width equal to the maximum cluster width. Hence, for the equivalent rectangular cubes, \( V = D \times E^2 \), where \( V \) is the tetrahedral volume, which is also the equivalent cube’s volume, \( D \) is the maximum cluster width that forms the cube’s width, and \( E^2 \) is the cross-sectional area of the cube.

We used step-wise polynomial fits to model the relationship between the uncertainty volume and the fiducial shift distance, and the maximum dipole cluster width and the fiducial distance.

**Random Fiducial Mismarkings**

Creating random combinations of fiducial shifts is another approach to analyze the effects of fiducial mismarkings. We generated 100 electrode location datasets for every 1-mm increment of the random fiducial mismarking combinations, producing 2000 datasets (100 datasets/increment \( \times \) 20 increments = total 2000 datasets). Each one-hundred mismarkings resided on a circular path with a fixed radius (\( 1 \leq r \leq 20 \) mm) away from the fiducial baseline locations. We ran DIPFIT using each dataset and compared the results with the outcomes of the systematic fiducial shifts.

**Results**

Shifting LPA, Nz, and RPA in different increments and directions resulted in dipole locations that had curvilinear paths in different planes. Each shift direction created changes in similar directions for the dipoles of the three ICs but with different magnitudes (Figure 3.2).

The uncertainty volume increased quadratically as a function of the magnitude of the fiducial shifts.
Figure 3.2: Fiducial mislocations and their corresponding estimated dipoles for A. Left preauricular (LPA), B. Nasion (Nz) and C. Right preauricular (RPA) for three independent components (ICs). Dipoles are located at the anterior cingulate (A.Cing.), the primary somatosensory cortex (Somat.) and at the premotor cortex (Premot.). Lighter colors show positive direction for the fiducial shifts. For each fiducial, resultant dipoles are plotted in sagittal, frontal and top views.

\( r^2 = 0.920, \) Figure 3.3A). The uncertainty volume was \(~0.06 \text{ cm}^3\) for fiducial shifts up to 0.5 cm for every IC. For fiducial shifts up to 1.3 cm, all three ICs showed similar increases in uncertainty volumes to \( 0.5 \text{ cm}^3\). With fiducial shifts >1.3 cm, the uncertainty volume of all ICs, including the three ICs of interest, began to separate from one another. At a 2 cm shift in fiducials, the average uncertainty volume was \( >2 \text{ cm}^3\).

The maximum dipole cluster width for an IC had a linear relationship with the fiducial shifts \( (r^2 = 0.79, \) Figure 3.3B). The average maximum cluster width exceeded 1 cm for shifts greater than 0.5 cm.
Figure 3.3: **A.** Relationship between the uncertainty volume and fiducial shifts. ICs in Figure 3.2 are drawn in color: anterior cingulate (ant. cingulate), somatosensory cortex (somatosens.) and premotor cortex (premotor). Tetrahedrons represent dipole uncertainty at the anterior cingulate. Pink cubes has the same volume as the tetrahedral volumes, with the same width as the maximum cluster width (D). The green dashed line quadratically models the uncertainty volume as a function of the fiducial shift. **B.** Maximum cluster width (D) of the dipoles estimated from the equidistant fiducial shifts. The green dashed line relates the maximum cluster width to the fiducial shift.

cm and was \( \sim 4 \) cm for fiducial shifts of 2 cm. Some of the cluster widths of different ICs started to deviate from the linear fit for the fiducial shifts >0.5 cm, although, we did not observe similar trend.
for the uncertainty volumes. The dipole located in the primary somatosensory cortex had larger maximum distances than the ICs in the anterior cingulate and premotor cortex for fiducial shifts greater than 1.3 cm.

Comparing results of the random fiducial mismarking combinations and the tetrahedral volumes from the systematic fiducial shifts, we found that 1) For each 1-mm increment, no random combination of the fiducial shifts (out of 100) could cancel out the effects of the shifts and result in a dipole that could make the uncertainty volume or cluster width smaller, and 2) >95% of the dipole estimations with random fiducial mismarking combinations at each 1-mm increment were in a close proximity of the corresponding tetrahedral surface (within \( \sim 20\% \) of the maximum cluster width).

Discussion and Conclusion

This study revealed the relationship of fiducial mismarkings during electrode digitization on the subsequent uncertainty of dipole location estimation. We found that shifts of a single fiducial location up to 0.5 cm resulted in an uncertainty volume <0.06 cm\(^3\) and a maximum distance <1 cm. When fiducial shifts were greater than 1.3 cm, dipole location uncertainty increased to >1 cm\(^3\) and the maximum distance increased to >2 cm.

One interesting finding was that the largest maximum distances, among the three ICs of interest, occurred in the primary somatosensory cortex, which is an area frequently discussed in the EEG studies related to walking [8,9]. A previous study found that tangential sources near the boundary of the cortex were more sensitive to electrode location errors, which could explain the larger maximum distances for the dipole at the primary somatosensory cortex compared to a dipole deeper within the cortex such as the anterior cingulate [5].

Another interesting finding was that the linear fiducial shifts mapped to curvilinear dipole paths in
different planes, which allowed the use of superposition to estimate dipole uncertainties created from fiducial mismarking combinations. Fiducial mismarking combinations are less likely to cancel each other, since the mismarking directions map to dipole paths in different planes. Fiducial mismarking combinations also created dipole spreads close to the uncertainty volumes that the systematic fiducial shifts had predicted.

Dipole cluster width was more sensitive to the fiducial shifts compared to volumetric uncertainty. Dipole cluster width had a steep linear relation with the fiducial shifts and mismarking the fiducials could change the location of a dipole as much as twice the fiducial mismarking shift. A recent study found that the reliability of a widely-used electromagnetic electrode digitizing system was $\sim 0.8$ cm [10]. Hence, even with perfect markings of the fiducials, there will still be up to 1.5 cm (17% of the head radius) uncertainty for the estimated dipole locations, just as a result of the reliability of the digitizing device.

Limitations of this study were that we did not examine every combination of the fiducial mismarkings and that we only co-registered the fiducials with the MNI head model for warping the electrode locations to the head model. While examining every combination of fiducial mismarkings was not practical, an advantage of analyzing random combinations of the fiducial mismarkings with the same distance from the baseline was that this approach could be used to estimate the mismarking effects for multiple digitizing systems if the digitizing reliability is known. While we could have co-registered more electrodes to the MNI head model, this would lose the individual characteristics of the digitized head. Alternatively, EEGLAB’s NFT toolbox enables warping of the MNI head model to all of the digitized electrode locations [11], but coupling NFT with 240 incremental electrode-location datasets was beyond our computational capacity.

Based on our results, we recommend using a digitizing system with measurement errors less than 0.5 cm and marking the fiducials within 0.5 cm of the actual fiducial to avoid errors greater than
1.5 cm in dipole location. Future work will compare the reliability of different digitizing systems to determine which digitizing systems have measurement errors less than the recommended 0.5 cm. While digitizing the actual locations of the EEG electrodes should provide greater dipole specificity, this study showed that small fiducial mismarkings could result in large dipole location uncertainty. To reduce dipole location uncertainty, care should be taken to minimize the cumulative potential errors from the user mismarking fiducial locations during the digitization process and the measurement errors from the digitization system.

References


CHAPTER 4: DIFFERENTIAL THETA-BAND SIGNATURES OF THE ANTERIOR CINGULATE AND MOTOR CORТИCES DURING SEATED LOCOMOTOR PERTURBATIONS

Introduction

Perturbing locomotion often produces error-driven adaptation where subjects adjust their locomotor patterns to reduce errors, but these adjustments revert to the unperturbed patterns after the perturbations are removed (i.e. wash-out) [1]. When subjects are re-exposed to the same perturbations or exposed to new perturbations, they adapt faster and may also modify unperturbed locomotor patterns [2]. However, these modifications may not transfer across lower limbs according to a split-belt walking study [3]. Despite the wash-out often seen with error-driven adaptation, split-crank cycling and split-belt walking can result in retained post-perturbation modifications if the modifications were not the direct task goals [4, 5]. For example, after cycling with different crank angles, subjects had perturbation-specific muscle activation patterns which did not wash-out post-perturbation [4]. These locomotor behaviors indicate that perturbations indeed modify locomotor responses beyond the perturbation period and that tuning perturbation features could modulate locomotor responses. Determining motor and cortical responses to different perturbations during a variety of locomotor tasks, beyond just walking, could greatly improve the understanding of locomotor adaptation processes.

Advancements in brain imaging technologies such as high-density electroencephalography (EEG), functional magnetic resonance imaging, and positron emission tomography have helped researchers

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identify supra-spinal correlates of locomotion [6–8]. The anterior cingulate cortex theta (3-8 Hz) power increases significantly during double-support in walking and during extension-onset in recumbent stepping [9, 10]. These studies suggest that the anterior cingulate activity may be monitoring more demanding locomotion phases. The supplementary motor area (SMA) has similar theta power fluctuations during walking, cycling, and recumbent stepping [6, 10, 11]. In general, the SMA and the motor cortex exhibit substantial alpha-beta (8-30 Hz) fluctuations during walking, with decreased alpha-beta power indicating active processing in the motor cortex [12].

The anterior cingulate cortex and SMA also both strongly respond to perturbations during walking and standing. Previous studies on split-belt walking, perturbed beam walking, walking over obstacles, and perturbed stepping reported perturbation-elicited activity of anterior cingulate, SMA, or in both areas [8,9,13–16]. The anterior cingulate cortex (or the equivalent mid-prefrontal cortex in the EEG channel studies) activity is often associated with error monitoring or motor learning, while SMA activity is associated with sensorimotor integration [15,17,18]. If the anterior cingulate cortex has an error monitoring function, we expect that the activity would scale with the error size [19]. If the anterior cingulate has a role in motor learning, we expect the activity would decrease with more perturbation experience. Previous studies on walking over obstacles and perturbed standing did not observe decreased anterior cingulate activity with more experience [13, 20]. On the other hand, a recent study reported scaling of central midline EEG signals at the Cz electrode with balance performance during a perturbed standing task [21]. However, all three studies had insufficient spatial resolution to determine confidently whether the electrocortical dynamics were from a single functional cortical area.

The purpose of this study was to determine the electrocortical signatures of motor responses to perturbations during a seated locomotor task. Adding perturbations during seated locomotor tasks such as recumbent stepping, which likely shares neural control with walking [22, 23] could provide an alternative option for gait rehabilitation since subjects do not need to maintain their balance.
Using our motorized recumbent stepper [24], we applied discrete mechanical perturbations during each stride and also had intermittent no-perturbation "catch" strides. The catch strides could probe whether subjects were updating anticipatory motor control strategies.

We had four hypotheses. The first hypothesis was that perturbations would initially create motor errors and increase anterior cingulate theta power near the perturbation event. As subjects gained more experience with perturbations, motor errors, and anterior cingulate theta power would decrease. The second hypothesis was that motor errors during the no-perturbation catch strides would increase the more subjects expected to encounter perturbations and that anterior cingulate spectral fluctuations would decrease in the later catches. The third hypothesis was that mid-extension perturbations, when the limbs were moving the fastest, would produce more significant errors and anterior cingulate theta power than extension-onset perturbations. We also expected to identify activity of the left and right motor cortices [10, 11] and hypothesized that spectral power fluctuations of the left and right motor cortices in response to the perturbations would be lateralized.

Methods

Subjects (n=17, 11 females, age 25 ± 4.9 years) performed perturbed arm-leg stepping on a one degree-of-freedom recumbent stepper (TRS 4000; NuStep, Inc., Ann Arbor, MI) integrated with a servomotor (Kollmorgen, Radford, VA), described in [24] (Figure 4.1a). The mechanically coupled left handle and right pedal move together out of phase with the mechanically coupled right handle and left pedal. As such, subjects could use any combination of their arms and legs to drive the stepper.
Figure 4.1: **Recumbent stepper and perturbations.**

- **a.** The recumbent stepper is a one-degree-of-freedom arm-leg exercise device.
- **b.** Perturbations were applied either at the extension-onset or mid-extension of the targeted leg. Perturbations were increased stepping resistance for 200 milliseconds.
- **c.** The experiment included four tasks. Each task had three ordered blocks, pre, perturbed stepping, and post. The perturbed stepping block also included random no-perturbation catch strides.

**Experiment procedure and motor errors**

The Institutional Review Board of the University of Central Florida approved the protocol and consent form, and the study was conducted per the principles stated in the Declaration of Helsinki. All subjects gave their written informed consent before starting the experiment. Subjects were all right-handed, based on the hand they would use to pick an object from the floor. Subjects
self-reported no prior neurological or musculoskeletal problems in the past two years before the data collection date.

We recorded EEG using a 128-electrode EEG system (ActiveTwo, BioSemi B.V., Amsterdam, the Netherlands). After placing the EEG cap on the subject’s head according to the BioSemi guidelines, we digitized the electrodes and fiducial locations using an infrared 3D scanner (Structure Sensor, Occipital Inc., Boulder, CO). We ensured that the resistance between the scalp and each electrode was <20 Ohms, indicating good contact between the electrodes and the scalp. We restrained cable movement using a cable holder behind the subject’s head and instructed subjects to keep their head steady to reduce EEG cable sway artifacts [25]. We strapped subjects’ feet to the pedals after they sat on the stepper seat. We also adjusted the handles to ensure that subjects were comfortable using the handles to drive the stepper.

The stepper’s servomotor perturbed the stepping motion with brief 200 ms increases in resistance at either the onset or middle of extension of the target (left or right) leg during the stepping stride (Figure 4.1b). The increased resistance magnitude during a perturbation required 3x the torque to maintain the stepping pace of 60 steps per minute. In total, there were four perturbation types (left/right leg * mid-extension/extension onset). A pacing cue equal to 60 steps per minute (=30 strides per minute) was provided on a visual display to help subjects maintain similar stepping speeds during and across tasks.

EEG was recorded at 512 Hz using the BioSemi software program, and the stepping kinematics were recorded using the servomotor’s encoder at 100 Hz in the stepper program. When the stepper program began and ended, a trigger signal was sent to start and stop the EEG recording to synchronize the data.
Figure 4.2: **Motor error metrics.** Subjects were instructed to match the pace and step smoothly. 

**a. Stepping duration (temporal)**

Stepping duration (i.e., temporal) error is the difference between stride duration and the two-second pacing cue.

**b. Stepping position (spatial)**

Stepping position (i.e., spatial) error is the maximum difference between the time-normalized stepping profile and the averaged *pre* profile.

**Data collection**

The data collection began with two minutes of quiet sitting, during which the pacing cues were shown as EEG was recorded. After completing this quiet sitting portion, subjects completed four 10-minute perturbed stepping tasks in a pseudo-randomized order. Each task only included one perturbation type. For each task, there were three ordered blocks: 1) **pre**: two minutes of unperturbed stepping, 2) **perturbed stepping**: six minutes of a single perturbation timing, and 3) **post**: two minutes of unperturbed stepping (Figure 4.1c). There were no pauses between blocks. In addition to perturbed strides, the perturbed stepping block included random one-in-five “catch” strides where no perturbation was applied. In this paper, we use pre and pre-perturbation and post and post-perturbation interchangeably. There was two minutes of quiet sitting at the end of the data collection.
Before starting each task, we instructed subjects to A) step smoothly as if they were walking, B) use both their arms and legs to drive the stepping motion, and C) follow the pacing cues that were projected in front of them (Figure 4.1a). We did not instruct subjects on how to follow the pacing cues as there are several options, such as having a leg be at full extension when the rectangle on the same side as the leg was black. Subjects also received no explicit feedback on whether they were stepping faster or slower than the pacing cue. Subjects were given at least two minutes of practice with the pacing cues before starting the data collection.

*Stride events*

After importing stepping data into MATLAB (R2018b, MathWorks Inc, Natick, MA), we separated each task into blocks and strides. We defined the strides as the time from one extension-onset of the perturbed leg to the next extension-onset of the perturbed leg. We excluded any incomplete strides. For each stride, we identified the following events: perturbed-step extension onset, perturbation (start time), recovery-step extension onset, and the end of the stride. We artificially added perturbation events to the unperturbed strides (i.e., pre, post, and catch strides), equal to the average latency of the perturbation events.

*Motor errors*

We quantified a temporal (pacing) error and a spatial (stepping) error, from the stepping kinematics (Figure 4.2). In our tasks, subjects should have completed a stride in two seconds based on the 60 steps-per-minute pacing cues. We defined temporal error as the stepping duration error, which was the difference between each stride duration and the two seconds (Figure 4.2a). Since we instructed subjects to step smoothly, we expected the stepping profiles to be smooth and rhythmic during the pre-perturbation block. We defined spatial error as a stepping position error, which was the
maximum difference between the time-normalized stepper position profile during each stride and the averaged pre-perturbation stepping profile (Figure 4.2b). We used the servomotor encoder data to quantify the angular stepping position around the stepper’s common rotating axis (Figure 4.1a).

**EEG processing**

EEG data were analyzed in MATLAB (R2018b, MathWorks Inc, Natick, MA) using a customized pipeline based on EEGLAB (version 2019.0) functions [26] (Figure 4.3). We used a high-pass filter at 1 Hz and a 60 Hz line-noise filter (CleanLine) to minimally clean the raw data [27, 28]. We imported stride events from the synchronized stepping data and concatenated data from all tasks into a single file. We then used a template correlation rejection method to identify and exclude channels with large cyclic artifacts [29].

![EEG post-processing workflow](image)

Figure 4.3: **EEG post-processing workflow with a novel step-wise algorithmic parameter sweeping noise rejection process.** Shaded blocks indicate inputs or outputs. Thick lined blocks highlight the novel step-wise rejection approach.

We developed and used a novel step-wise channel and frame rejection algorithm to reject channels and data frames that still contained considerable noise (Figure 4.3). We removed the researcher’s need to set single thresholds for the channel and frame rejection steps. Instead, the step-wise algorithm identified a suite of thresholds, from lenient to conservative, that created 32 separate datasets with different rejection levels for each participant (8 steps for channel rejection * 4 steps...
for data-frame rejection = 32 datasets). Channel rejection metrics were the signal range, standard deviation, kurtosis, and correlation to the other channels. Frame rejection involved finding periods of the EEG data with a significantly higher signal variability than the overall median of signal variability. While the number of the rejected channels and frames varied for each participant and increment, we set the rejection thresholds such that the most conservative increment always retained > 85 channels and > 80% of data.

We used independent component analysis (ICA), the dipolar source estimation technique (DIPFIT), and a multi-variate source classifier (ICLabel) on each step-wise dataset to identify and locate the sources that contributed to the EEG signals. We specifically used the adaptive mixture independent component analysis (AMICA) to separate the EEG into temporally independent components [30]. To select the best increment from the 32 step-wise datasets, we first estimated the source locations for each dataset’s independent components using EEGLAB’s DIPFIT version 3.0. We then excluded any source located outside the brain or with the residual variance > 15%. We then used EEGLAB’s ICLabel toolbox to classify the source types as “brain” or “non-brain” [31] and selected the step-wise dataset with the topmost “brain” sources as the representative dataset for each subject. We visually checked the results of the ICLabel for the selected dataset to confirm the classification of the sources as “brain” (or “non-brain”).

We then clustered the sources across all subjects based on the source location, power spectrum, and scalp map. We divided the power spectrum and scalp map into ten bins. The binned power spectrum was from 3 to 25 Hz. The Laplacian of the scalp map was used for clustering [32]. We developed and used a novel optimal k-means approach to determine the number of clusters from a range of possible numbers of clusters provided to the algorithm (here, from 15 to 30 clusters). The optimal k-means approach uses MATLAB’s “evalcluster” function to find the specific number of clusters that maximize the similarity of the sources within each cluster. We kept and analyzed only the clusters that contained components from more than 70% of the subjects. If a subject had
multiple sources in a cluster, we only kept the source with the largest channel data variance. We identified Brodmann Areas and cortical cortices of the sources and cluster centroids using Talairach coordinates and talairach.org [33, 34].

We computed the time-frequency spectral power of each source in the cluster across the stride epochs, known as event-related spectral perturbations (ERSP) [35]. For hypotheses 1, 2, and 4, each epoch was a stride, and for hypothesis 3, each epoch was –400 ms to +400 ms of the perturbation event. We padded the epochs by 700 ms to avoid possible edge effects. Next, we baseline-normalized the spectral power based on the pre-perturbation block’s average spectral power and computed the ensembled average ERSPs across subjects. We determined the significant event-related synchronization and event-related desynchronization across the ERSPs using EEGLAB’s bootstrapping method with alpha set at 0.05 [36]. ERSP images only show significant spectral fluctuations.

Statistical analysis

Identification of motor responses

We tested the temporal and spatial errors to determine the error behavior. For each subject, we divided their strides into 20% batches in the pre and post blocks and 10% batches in the perturbed-stepping block [37]. We compared the average of the first and last batches of perturbed strides, catch strides, and post-perturbations strides, as well as the last batch of pre-perturbation strides using repeated-measure analysis of variance (rANOVA) for each error and task. If the rANOVA was significant, we performed a priori Fisher’s Least Significant Difference (LSD) tests for the following pairs: 1) late pre vs. late post (for sustained post modifications and wash-out), 2) early vs. late post (for wash-out), 3) early vs. late catch (for adaptation), and 4) early vs. late perturbed (for
Figure 4.4: **Temporal and spatial errors during perturbed stepping.** The *perturbed-stepping* block includes both perturbed strides (green) and one-in-five random catch strides (purple). For *perturbed-stepping*, the 10 circles are the averages of 10% batches. For *pre* and *post*, the 5 circles are the averages of 20% batches. * indicates significant post-hoc LSD tests. **a.** Temporal errors were different between perturbed and catch strides (~50 ms vs ~200 ms) and returned to the *pre* levels during the *post* block. **b.** Spatial error was greater for the perturbed strides than catch strides, and did not return to *pre* levels during *post*.

adaptation). The significance level for all statistical tests was 0.05.

**Tests for hypotheses 1 and 2: adaptation of motor and cortical responses**

We compared motor errors and electrocortical dynamics between the early (first 33% of the strides) and late (last 33% of the strides) in the perturbed stepping block. Here, we used 33% of the strides as early or late to retain at least 10 strides per subject (total perturbed strides≈140-150, total catch strides≈30-40) for EEG group-level analyses [35, 36]. We tested motor errors for the perturbed and catch strides separately and used rANOVA with three factors: 1) *adaptation* with two levels: early and late, 2) *task side* with two levels: left and right, and 3) *perturbation timing* also with two
levels: mid-extension and extension-onset. We performed post-hoc Student paired t-tests only if the adaptation had a significant main effect because adaptation was the only factor pertinent to our first two hypotheses. To compare the electrocortical responses between early and late perturbed strides and early and late catch strides, we computed the spectral fluctuations and averaged the spectral powers to derive the theta-band (3-8 Hz) ERSP waveform [36, 38]. We compared the early and late theta-band average ERSP waveforms using bootstrapped paired t-tests and false discovery rate corrections for multiple comparisons with EEGLAB’s “statcond” and “fdr” functions. We also determined meaningful spectral-power increases or decreases of the theta-band by determining when the power confidence interval was greater or less than zero. We excluded other frequency bands because preliminary analyses showed the main spectral fluctuations were limited to theta. The significance level for all statistical tests was 0.05.

Tests for hypothesis 3: effect of perturbation timing

We included all perturbed strides to quantify possible motor and electrocortical differences between perturbation timings, i.e., mid-extension and extension onset. For each motor error, we used rANOVA with two factors: 1) perturbation timing with two levels: mid-extension and extension-onset, and 2) task side also with two levels: left and right. We only performed a post-hoc Student paired t-test between the same side tasks if there was a significant perturbation timing effect. We compared the ERSPs centered around the perturbation event for left-side tasks (i.e., left mid-extension and left extension-onset) and right-side tasks separately. Similar to the tests for hypotheses 1 and 2, we used bootstrapped paired t-tests with corrections for multiple comparisons to compare the theta-band average ERSP between the tasks and determined meaningful spectral-power increases or decreases when the power confidence interval cleared zero. All statistical tests had 0.05 significance level.
Tests for hypothesis 4, motor cortex lateralization

We compared spectral fluctuations of the cortical clusters during the left and right-side tasks to investigate contralateral and specialized lateralization of the motor cortex. Hemispheric activity that corresponds to contralateral limb movements is contralateral lateralization whereas hemispheric activity that corresponds with ipsilateral limb movements is specialized lateralization [39]. All perturbed and catch strides were included in this analysis.

Results

Motor error responses and cortical clusters

Temporal (pacing) and spatial (stepping) errors did not decrease with more exposure to perturbations, and spatial errors did not wash-out (Figure 4.4). Perturbed-stride temporal errors were ~50 ms but catch-stride temporal errors were ~200 ms (Figure 4.3a). The rANOVAs indicated significant temporal error differences in each task ($F'(6,96)>40, p'<0.0005$). Post-hoc LSD showed a slight temporal error increase during left and right extension-onset perturbed strides and a temporal error decrease during right extension-onset strides post-perturbation. Spatial errors of perturbed strides were steady, ~12° for mid-extension and ~16° for the extension-onset (Figure 4.3b). During the mid-extension tasks, catch-stride spatial errors seemed a continuation of pre-perturbation errors at ~5° but trended to ~10° by the end of the catch strides. rANOVAs were significant for the spatial errors across all tasks ($F'(6,96)>29, p'<0.0005$). Spatial errors did not wash-out (i.e., return to pre) during post-perturbation (post-hoc LSD p’s<0.05). However, post-perturbation spatial errors in the right-side tasks decreased from the first to last batch (LSD p’s<0.05).

The optimal k-means identified five cortical clusters (Figure 4.5). We focused on three clusters
located at the anterior cingulate cortex (14 subjects), left SMA (13 subjects), and right SMA (13 subjects). Cluster locations were assigned to the nearest Brodmann areas based on the Talairach coordinates of the cluster centroid [34]. As the SMA and premotor cortex share Brodmann area 6, we further confirmed the SMA cluster locations from a previous fMRI and PET meta-analysis [40]. The left and right SMA were determined based on the cluster’s centroid location and the individual source locations.

Figure 4.5: **Locations of the electrocortical clusters.** Clusters with sources from > 70% of the subjects are shown. Only one source per subject was selected for each cluster during analysis. "% of all" indicates the percentage of all components in the Brodmann Area. a.u.: arbitrary unit

<table>
<thead>
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<th>no. of subjects</th>
<th>14</th>
<th>13</th>
<th>13</th>
<th>13</th>
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<td>no. of ICS</td>
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<td>29</td>
<td>33</td>
<td>24</td>
<td>24</td>
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<td>Centroid Talairach coordinates</td>
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<td>(-8.1, 2.3, 61.2)</td>
<td>(8.3, 15.5, 64.0)</td>
<td>(-15.3, -46.4, 53.2)</td>
<td>(38.8, -63.7, 42.7)</td>
</tr>
<tr>
<td>Brodmann area (% of all)</td>
<td>24, 32 (74%)</td>
<td>6 (83%)</td>
<td>6 (88%)</td>
<td>7 (88%)</td>
<td>7 (42%), 39 (33%), 40 (21%)</td>
</tr>
</tbody>
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Anterior cingulate theta-band adaptation occurred without motor error adaptation in perturbed strides

Motor errors of the perturbed strides did not decrease from early to late, but anterior cingulate theta-band spectral power decreased in the right-side tasks (Figure 4.6a-c). Neither adaptation nor task side had a significant effect on the perturbed strides (rANOVA, temporal adaptation: F(1,16)=0.74, p=0.789, temporal side: F(1,16)=0.74, p=0.789, spatial adaptation: F(1,16)=2.98, p=0.104, spatial side: F(1,16)=0.71, p=0.413). Mid-extension perturbed strides had significantly greater average temporal errors (71 vs 39 ms) but smaller spatial errors (12° vs 16°) than the extension-onset
perturbed strides (rANOVA, temporal: $F_{(1,16)}=461$, p<0.0005, spatial: $F_{(1,16)}=30.6$, p<0.0005). Perturbations elicited anterior cingulate theta synchronization during all tasks (Figure 4.6b). Theta spectral power decreased from early to late for right-side perturbed strides. The left-side perturbed strides, however, had similar and sometimes stronger theta synchronization during late strides than the early strides. The anterior cingulate cortex also showed theta desynchronization in the recovery steps (i.e., the unperturbed steps after perturbed steps), specifically for the early right-side and late left-side perturbations. The theta-band average ERSP bootstrap t-tests revealed that spectral power had a decreasing trend from early to late for the right-side tasks, which was significant for right mid-extension perturbations (Figure 4.6c). In the left-side perturbed strides, late synchronizations during the recovery steps were statistically different from the non-significant spectral fluctuations during early recovery steps.
Figure 4.6: **Motor errors and anterior cingulate ERSP** a. and d. Adaptation (early vs. late) was not a significant factor for motor errors. # indicates perturbation timing was a significant factor. Error bars indicate confidence interval (CI) b. Perturbations (i.e., the strong solid lines) elicited theta synchronization in anterior cingulate cortex. Right-side perturbations elicited weaker synchronization during the late perturbed strides. c. Average theta-band ERSP waveform shows increased power after the perturbations across the tasks. Late perturbations elicited less theta-band average power only on the right-side tasks. e. Early catches (narrow solid lines) elicited a theta synchronization in the anterior cingulate cortex. f. Average theta-band ERSP waveform shows late right extension-onset catch strides elicited significantly higher spectral power than the early catch. Shaded areas indicate CI. Black bars indicate significant difference between early and late. Colored bars indicate CI does not overlap with zero.
Motor errors did not increase from early to late catch strides, but early catch strides still elicited theta synchronization in the anterior cingulate cortex (Figure 4.6d-f). Similar to the perturbed strides, neither adaptation nor task side had a significant effect on the temporal or spatial motor errors (rANOVA, temporal-adaptation: F(1,16) = 3.76, p = 0.070, temporal-side: F(1,16) = 1.65, p = 0.217, spatial-adaptation: F(1,16) = 1.43, p = 0.25, spatial-side: F(1,16) = 0.70, p = 0.415). Mid-extension catch strides had significantly greater average temporal (250 vs 153 ms) and spatial errors (8° vs 6°) than the extension-onset catch strides (rANOVA, temporal: F(1,16) = 31.9, p < 0.0005, spatial: F(1,16) = 24.4, p < 0.0005). Early catch steps elicited anterior cingulate theta synchronization (Figure 4.6e). This synchronization occurred before completion of the catch-step extension for the mid-extension tasks but was near the start of the catch step for just the right extension-onset catch strides. The left mid-extension and right extension-onset elicited theta desynchronization in the recovery steps (Figure 4.6e-f). For right extension-onset, the recovery steps of the late catch strides elicited significantly greater spectral power than the early catch strides.

Group analyses including all perturbed strides revealed differential motor error and cortical responses based on perturbation timing (Figure 4.7). Comparing motor errors across all perturbed strides revealed that perturbation timing, and not the task side, was a significant factor (rANOVA, temporal timing: F(1,16) = 30.5, p < 0.0005, temporal side: F(1,16) = 2.24, p = 0.15, spatial timing: F(1,16) = 27.5, p < 0.0005, spatial side: F(1,16) = 0.25, p = 0.62) (Figure 4.7a-b). For each side, temporal error was significantly greater during mid-extension than extension-onset and the spatial error was smaller during mid-extension than extension onset (post-hoc paired t-test, temporal right: p < 0.0005, tempo-
Figure 4.7: **Motor errors and theta-band ERSP across perturbation timings.** a. and b. Task type (mid-extension vs. extension-onset) was a significant factor for motor errors. Error bars indicate confidence interval (CI). c. Extension-onset perturbations had greater anterior cingulate and ipsilateral theta-band ERSP before the perturbation event (the solid vertical line). The theta-band ERSP was not significantly different after the perturbation event, except for the left SMA ipsilateral task. Shaded area indicates confidence interval. Black bars indicate significant difference between perturbation timing. Colored bars indicate CI clearing zero.

rual left: p=0.001, spatial right: p<0.0005, spatial left: p=0.001). The extension-onset perturbations elicited a significant increase in theta-band ERSP before the perturbation event across the anterior cingulate and ipsilateral SMA (Figure 4.7c). The increased theta-band ERSP for mid-extension
perturbations was delayed, occurring after the perturbation event in the anterior cingulate and left SMA but was about -100 ms before the perturbation event in the right SMA. Only the ipsilateral left SMA showed greater extension-onset theta-band ERSP than mid-extension for more than 100 ms after the perturbation onset.

*Cortical lateralization and specialization*

The left and right SMAs demonstrated both contralateral and task-specific lateralization with respect to lower limb extension (Figure 4.8). The recovery-step desynchronization during the perturbed strides was most prominent in the right SMA for extension-onset tasks and in the left SMA for the mid-extension tasks, indicating presence of task-specific lateralization (Figure 4.8a, red rectangles). Mid-extension tasks also involved theta desynchronization before the perturbation event in both left and right SMAs (Figure 4.8a, red dashed rectangles). Similar recovery-step desynchronization was present in the catch strides but were limited to the ipsilateral SMA of the recovery-step leg during extension-onset tasks and to the right SMA for mid-extension tasks (Figure 4.8b, black and red rectangles). Strong theta synchronization only occurred in the right SMA for mid-extension catch strides just before the end of the catch step (Figure 4.8b, red dashed rectangles).

**Discussion**

We quantified motor and electrocortical responses to frequent mechanical perturbations during recumbent stepping to gain insight on the electrocortical dynamics of locomotor adaptation. We did not observe typical motor error adaptation. Temporal errors were consistently ~50ms of the desired pace during perturbed strides and returned to pre-perturbation levels in the post block. Spatial errors did not adapt (decrease) with more exposure to the perturbations and did not return to
Figure 4.8: Supplementary motor area (SMA) Lateralization. a. Theta desynchronization in the perturbed recovery step occurred in the right SMA for extension-onset and in the left SMA for the mid-extension (red squares). Only mid-extension perturbations elicited theta desynchronization before the perturbation event (red dashed rectangles). b. Mid-extension catch steps elicited theta synchronization before the end of limb extension (red dashed rectangles). Recovery-step theta desynchronization occurred contralaterally during extension-onset, but only occurred in the right SMA during the mid-extension. Red and black indicate specialized and contralateral lateralization respectively.

pre-perturbation levels in the post block. The lack of error-based adaptation behavior coupled with small temporal errors and sustained spatial errors in the post block are indicative of use-dependent learning [41]. Electrocortical sources in the anterior cingulate cortex and supplementary motor...
areas showed that perturbations elicited theta synchronization, as expected. Despite the lack of motor error adaptation, anterior cingulate theta synchronization showed a decreasing trend during late perturbed strides in the right-side tasks. Interestingly, theta-band ERSP during extension-onset tasks started before the perturbation event, resulting in greater theta synchronization in the anterior cingulate and SMAs preceding the perturbation event compared to mid-extension tasks. Motor cortex lateralization was mostly task-specific, where theta desynchronization occurred during the recovery-step in the right SMA for extension-onset tasks but in the left SMA for the mid-extension tasks. These results highlight that electrocortical and motor responses are not necessarily coupled and that perturbation features such as timing could be tuned to elicit greater involvement of specific brain areas.

The perturbed recumbent stepping protocol did not produce the typical error reduction and rapid wash-out associated with motor adaptation, but instead, revealed sustained errors during the post-perturbation block (Figure 4.4), suggesting use-dependent learning occurred. In preliminary analyses, we compared multiple definitions of early and late to determine the robustness of the lack of error-based adaptation in our study. Statistical tests consistently showed no significant difference between early and late, except when early was defined as just the first stride. When perturbations do not directly hinder achieving the task goal, use-dependent learning emerges more than error-based adaptation [41]. With use-dependent learning, motor behaviors are modified in the direction of perturbation and sustained longer after removing the perturbations. Here, we did not provide subjects with any visual feedback of their errors or task performance, so, matching the stepping pace with the pacing cues was the more explicit task goal. Because subjects matched the pacing cues well with temporal errors of ~50 ms, which might be imperceptible for active control adjustments [42], the perturbations did not hinder achieving the task goal. For the less explicit goal of stepping smoothly, however, spatial errors were sustained during perturbed strides and did not wash out during the 2-minute post-perturbation block. During split-crank cycling and split-belt walking
where changing muscle recruitment was not an explicit task goal, modified muscular activation patterns were sustained [4, 5]. More recently, a perturbed walking study using brief treadmill belt accelerations during push-off also reported use-dependent learning and sustained post-perturbation gait modifications [43]. Longer-lasting locomotor modifications are desirable for gait rehabilitation and warrant further development of use-dependent learning paradigms.

Perturbations during our seated locomotor task elicited significant anterior cingulate theta synchronization that also decreased with time (i.e. adapted) for right-side perturbations, providing new insights about the anterior cingulate role in error monitoring and motor learning. Previous studies have attributed anterior theta synchronization, or the analogous negative deflection in event-related potentials, to physical loss of balance or presence of a postural threat [15, 44]. Our results demonstrated that even without a potential loss of balance, mechanical perturbations during a seated locomotor task can elicit anterior cingulate activity. We also observed a trend of adaptation of anterior cingulate theta synchronization for the right-side tasks, which contrasts previous studies that did not observe changes in the anterior cingulate cortex with adaptation but acknowledged a lack of spatial resolution [13, 20]. Our approach likely had sufficient resolution [34, 45], but we observed a trend of anterior cingulate adaptation only in the right-side tasks. Despite consistent motor errors during catch strides, only early catch strides elicited theta synchronization, suggesting that the anterior cingulate perceived early catches as errors, which emphasizes that mechanical perturbations are crucial for anterior cingulate elicitation. Overall, the sustained anterior cingulate theta power across all tasks during perturbed stepping further supports that the anterior cingulate cortex has a role in error-monitoring. However, the theta-band adaptation trend during right-side perturbed stepping suggests that the anterior cingulate also has a role in locomotor learning.

Perturbation timing significantly influenced anterior cingulate theta-band power fluctuations (Figure 4.6), suggesting that tuning perturbation features can modify and stimulate anterior cingulate activity. The theta-band average ERSP for extension-onset perturbations was greater than mid-extension
perturbations. This difference may result from an additional intrinsic anterior cingulate theta synchronization that occurs during limb transitions in unperturbed gait, pedaling, and stepping [6, 7, 9, 10]. However, our results did not show significant anterior cingulate activity during pre and post-perturbations strides, partly because our analyses and ICA focused on identifying sources involved in perturbed stepping. The sustained anterior cingulate theta-band elicitation over the entire six minutes of left mid-extension perturbations demonstrates that specific perturbations could be tuned to enhance or extend cortical engagement.

We identified two close but distinct SMA clusters that exhibited specialized lateralization with both theta synchronization and desynchronization. We were able to identify distinct clusters in close proximity using our novel EEG noise rejection process that performs algorithmic parameter sweeping to estimate the most brain sources and an optimal k-means clustering algorithm to identify optimal cortical clusters (Figure 4.3). The left and right SMAs had clear differences in theta fluctuations (Figure 4.8), supporting that these SMA clusters were distinct and had specialized responses to the perturbations or motor errors. Theta synchronization occurred exclusively in the right SMA during mid-extension catch steps that had the largest temporal errors (~250 ms), suggesting that despite the lack of a physical perturbation, the right motor area theta synchronization was still sensitive to a motor error. The right motor and premotor cortices have been linked with monitoring temporal aspects of motor tasks [38, 39].

Interestingly, theta desynchronization occurred during the recovery step (i.e., the step following a perturbed step) in the right SMA during extension-onset perturbations and in the left SMA during mid-extension perturbations (Figure 4.8). Previous unperturbed gait studies showed significant theta desynchronization in sensorimotor cortices during mid-stance, but the significance of theta desynchronization specifically is not discussed [6, 7, 9, 10]. A recent study demonstrated that theta synchronization and desynchronization corresponds to negative and positive deflections in event-related potentials (ERP) of motor cortex, respectively [46]. As such, ERP studies provide
additional possible interpretations for observed theta synchronization and desynchronization in locomotor tasks. For example, a recent study on upper-limb visuomotor perturbations suggested that the presence (or absence) of negative and positive potentials during perturbations indicated different motor learning strategies [47], which aligns with our results.

Limitations of this study include attributing cortical and motor responses to lower-limb extension and focusing on EEG group-level analyses. We attributed the perturbations to the action of extending the lower-limb, i.e., left mid-extension perturbation means the perturbation occurred in the middle of extending the left-leg. Our stepping torque analysis (not reported here) and a previous study showed that during arms and legs recumbent stepping, subjects mainly relied on lower-limb extension for higher power demands [48]. Ensemble averaging across strides and subjects is necessary for EEG group-level analysis to reveal event-locked cortical fluctuations [35, 36]. Previous studies had >60 strides per subject for ensemble averaging, inherently increasing their statistical power [15, 38]. We had ~10 catch strides and ~50 perturbed strides per subject and yet were still able to observe distinct event-locked spectral fluctuations. Further single trial analysis may provide more insights into the inter-stride cortical variability [20, 49].

Mechanical perturbations are a robust way to elicit error-related cortical fluctuations and could be tuned to further enhance desired cortical activity. During a seated locomotor task, mechanical perturbations elicited anterior cingulate cortex activity, which decreased with more experience with the right-side perturbations. This supports that the anterior cingulate both monitors errors and learns from them [50]. The left and right SMA clusters demonstrated task-specific lateralization, suggesting that tuning perturbation features such as timing can elicit more desired cortical activity. The uncoupled anterior cingulate activity with motor errors and the specialized SMA fluctuations implicate that cortical feedback may be crucial for closed-loop rehabilitation because motor changes may not adequately reflect cortical dynamics.
References


CHAPTER 5: OLDER ADULTS USE FEWER MUSCLES TO OVERCOME PERTURBATIONS DURING A SEATED LOCOMOTOR TASK

Introduction

Muscle co-contraction, i.e., the concurrent activity of the agonist and antagonist muscles, is a common strategy when responding to motor perturbations and during increased uncertainty; this co-contraction usually decreases with the progression of adaptation and reduction of motor errors in response to the perturbations [1, 2]. Young adults seem to use co-contraction more strategically to adapt more rapidly to perturbations and improve their accuracy in upper limb reaching tasks [3, 4]. Young adults initially increase their muscular activity in response to postural balance challenges and split-belt walking [5, 6]. This muscular activity gradually decreases after adapting to the perturbations and reducing motor errors. However, young adults may not reduce their co-contraction as they adapt to standing or walking perturbations [6, 7].

Older adults often use more global co-contraction compared to young adults, presumably to resist perturbations [8–10]. During perturbed goal-directed reaching tasks, older adults did not reduce their motor errors or co-contraction as much as young adults [8, 11]. During postural and locomotor perturbations, older adults also used more co-contraction, indicating an increased effort to adapt to the perturbations [7, 9]. An undesirable consequence of the increased co-contraction during postural tasks is that it may reduce balance performance, particularly in older adults [12]. The increased co-contraction during balance tasks and walking in older adults seems to be an age-specific strategy, which is not due to a lack of sensory acuity and might be insufficient to respond to losses of balance [10, 12–15]. Nonetheless, older adults can improve their walking and balance performance and reduce their co-contraction as they gain more experience with the perturbations during postural
tasks and walking [16, 17]. These reductions in co-contraction may not translate to improved balance or walking metrics, however [18–20].

Use-dependent learning produces a prolonged adaptation of movements that do not wash out in a few strides after removing the perturbations [21]. During use-dependent learning, perturbations do not directly hinder the completion of the task such that reducing motor task errors is necessarily advantageous. For example, applying brief belt accelerations at toe-off of each leg on a split-belt treadmill would not challenge balance such that subjects learned to increase push off in response to perturbations and retained the stronger push offs even after the perturbations were removed [22]. We recently showed that perturbing recumbent stepping using brief increases in resistance did not produce classic error-based adaptation but rather, resulted in features of use-dependent learning in young adults [23]. The brief resistive perturbations did not hinder the most explicit task goal of following a pacing cue. As such, subjects modified their stepping patterns without reducing temporal or spatial errors, and these modified patterns were sustained even after removing the perturbations and stepping without perturbations for 2 minutes [23]. During perturbed cycling using a split-crank that altered the relative phasing of the pedaling legs, subjects modified their muscle activity patterns and retain those patterns [24]. The potential for shaping muscle activity, co-contraction, and motor behavior using use-dependent learning tasks has not been explored much, particularly during seated locomotor tasks.

The purpose of this study was to compare motor behavioral and muscular responses to perturbations during recumbent stepping in young and older adults. We expected older adults to respond to perturbation similarly to young adults [23] in that the older adults would not show error-based adaptations. We hypothesized that perturbations would increase muscle recruitment and be sustained after the perturbations were removed, consistent with use-dependent learning in young and older adults. Additionally, we hypothesized older adults would exhibit more muscle co-contraction compared to young adults.
We used our robotic recumbent stepper to perturb young and older adults during recumbent stepping by briefly increasing the stepping resistance. Subjects completed four perturbed stepping tasks; each task involved a single perturbation that occurred at extension-onset or mid-extension of the left or right leg. We instructed subjects to use both their arms and legs, but subjects could drive the stepper with only one limb as the recumbent stepper has only one-degree-of-freedom. We recorded the stepping kinematics and subject’s EMG from twelve muscles and quantified motor errors, mean EMG, and the co-contraction index.

**Methods**

Seventeen young adults (11 females, age 25 ± 4.9 years) and 11 older adults (4 females, age 68 ± 3.6 years) participated in the study. Subjects were all right-handed based on which hand they would use to pick up an object from the floor. They self-reported no neurological impairments, no problems with their gait, no history of falls, and no broken bones for two years before the data collection. Each participant also met the inclusion criteria based on four questionnaires to ensure they could safely complete the experiment: 1- Short performance battery (9/12) [25], 2- Berg balance scale examination (50/56) [26], 3- Mini mental-state examination (25/30) [27], and 4- CHAMPS physical activity [28]. The Institutional Review Board of the University of Central Florida approved the study, and subjects gave their written informed consent before starting the experiment.

**Hardware**

We used a recumbent stepper integrated with a servomotor [29] to introduce brief perturbations in the form of added resistance during stepping (Figure 5.1a). The stepper (TRS 4000; NuStep, Inc., Ann Arbor, MI) was mechanically coupled in a way such that the contralateral arm and leg
would extend together. We used the servomotor’s position sensor (Kollmorgen, Radford, VA) to record the stepper’s kinematics at 100Hz. Perturbations briefly increased stepping resistance for 200 milliseconds. The magnitude of the resistance required 3x torque to drive the stepper at 60 steps per minute. Perturbations were applied once the targeted leg was at the extension-onset or the mid-extension (Figure 5.1b).

We used twelve wireless electromyography (EMG) sensors (Trigno, Delsys, Natick, MA) to record muscular activity at ~1.1 kHz from the tibialis anterior, soleus, rectus femoris, semitendinosus, anterior deltoid, and posterior deltoid on both the left and right limbs. After locating the sensor position according to the SENIAM guidelines [30], we abraded and cleaned the skin and attached the sensors using the Delsys double-sided adhesive patches. Data streams of the EMG and stepper systems were synchronized using a trigger signal sent from the stepper controller to the EMG controller to start and stop recording simultaneously. We imported and preprocessed the stepper data in MATLAB (R2018b, MathWorks Inc, Natick, MA). We completed all EMG processing, as well as stepping motor error quantification in Python 3.8.2, using Numpy 1.19 [31], Scipy 1.6 [32], Pandas 1.2 [33], and Matplotlib 3.3 [34].

Protocol

Data collection started with two minutes of quiet sitting, was followed by four 10-minute stepping tasks, and ended with another two minutes of quiet sitting. Each stepping task only included one type of perturbation, i.e., two perturbation windows (extension-onset or the mid-extension) x two legs = four perturbation type. The order of the perturbed trials was pseudorandomized. Each perturbed stepping task included three different blocks (Figure 5.1c): 1) pre: two minutes of unperturbed stepping at the start of each trial. 2) perturbed stepping: six minutes of perturbed strides with a single perturbation type. 3) post: two minutes of unperturbed stepping immediately
Figure 5.1: Schematic of the robotic recumbent stepper, perturbations types, and stepping blocks. 

a. The robotic recumbent stepper is a one-degree-of-freedom stepping device with an integrated servomotor. The handles and pedals are mechanically coupled such that any limb can drive the stepping motion and move all of the other limbs. A pacing cue of alternating black and white rectangles that were 180 degrees out of phase with another was projected on a screen in front of the subject. 

b. Perturbations were brief increases in stepping resistance in the extension-onset or mid-extension of each stride (shaded light green vertical rectangle).

c. Each task block consisted of six minutes of perturbed stepping padded by two minutes of unperturbed stepping at the beginning and end of the task. Random catch strides did not include a perturbation.

after the end of the perturbed stepping period. The perturbed stepping block also included a random “catch” stride in every five perturbed strides, which did not apply a perturbation. We use pre and pre-perturbation interchangeably and also use post and post-perturbation interchangeably.

We strapped the subject’s feet on the pedals, adjusted the seat position, and moved the handles to ensure subjects would not lock their knees and could drive the stepper with the handles easily. Before each task, we instructed the subjects to A) step smoothly as if they were walking, B) use both their arms and legs to drive the stepper, and C) follow the pacing cues that were projected in front of them (Figure 5.1). Pacing cues were set at 60 steps per minute to match older adults’ average walking pace [35] and were projected as two reciprocating black and white rectangles (Figure 5.1).

We did not provide any instruction on how to interpret the pacing cues. Subjects were given at least two minutes of training to become familiar with the pacing cues before starting the data collection.
Stepping preprocessing and stride events

We separated each task into blocks and strides after importing the stepping data into MATLAB. We defined a stride as the time from one extension-onset of the perturbed leg to the next extension-onset of the perturbed leg for each task. For each stride, we identified the following events: perturbed-leg extension onset, perturbation (start time), unperturbed-leg extension onset, and the end of the stride. We artificially inserted perturbation events to the unperturbed strides (i.e., pre, post, and catch strides), at the average latency the perturbation events during the perturbed strides. We excluded any incomplete strides, which were strides that did not include all the events.

Motor Errors

We quantified two motor error metrics, one temporal and one spatial, from the stepping kinematics. Based on the pacing cues at 60 steps-per-minute, subjects should have completed each stride in two seconds. We defined temporal error as the stride duration error, which was the difference between each stride duration and the two seconds (Figure 5.2a). Because we instructed subjects to step smoothly, we expected the stepping profiles to be smooth and rhythmic during the pre-perturbation block. We defined spatial error as a stepping position error, i.e., the maximum difference of the time-normalized position profile during each stride from the averaged pre-perturbation stepping profile (Figure 5.2b).

EMG processing

We imported and analyzed the EMG data in the Python environment using a custom processing pipeline based on Banks et al. [36]. We resampled the EMG data to 1 kHz, band-pass filtered between 30 and 200 Hz, rectified, and low-pass filtered at 20 Hz to obtain the EMG linear envelopes.
Filters were designed using the 6th-order Butterworth algorithm. We chose 20 Hz as the low-pass threshold to capture EMG fluctuations in response to our 200-ms perturbations [37]. We then epoched and time-normalized the EMG data based on the stepping events for each stride. Finally, we normalized each muscle’s linear envelope to the overall average of the muscle’s linear envelope across all tasks.

Table 5.1: Agonist muscles to drive the stepper for the left- and right-side tasks. L = left. R = right.

<table>
<thead>
<tr>
<th>Muscle pairs</th>
<th>Agonist muscle</th>
<th>Left-side tasks</th>
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<td></td>
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<td>Perturbed step</td>
<td>Recovery step</td>
<td>Perturbed step</td>
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<tr>
<td>L. anterior deltoid – L. posterior</td>
<td>L. posterior</td>
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We used the ‘fixed’ approach to quantify co-contraction [36]. We assumed that the agonist was the muscle that could drive the stepper without the activity of the other muscles. During the step that involved left leg extension, the left soleus, left rectus femoris, left posterior deltoid, right tibialis anterior, right semitendinosus, and right anterior deltoit act as functional agonists. The agonist muscles of the muscle pairs for each step are summarized in Table 1. The fixed co-contraction index (CI) is calculated using the following equation:

\[ CI = \frac{2 \times I_{\text{antagonist}}}{I_{\text{agonist}} + I_{\text{antagonist}}} \]
Here, $I_{\text{antagonist}}$ and $I_{\text{agonist}}$ are the integrals of the EMG linear envelopes over each step. Because of the stepper’s inherent redundancy, subjects may use a subset of agonists to drive the stepper and use a few antagonist muscles to control stepping. CI is usually expected to remain $<1$, so the net activity of the muscle pair can drive the limb in the designated stepping direction. However, in our study, CI might become $>1$ if the designated antagonist helps to control stepping while the agonist is not involved in driving the stepper. CI $<1$ means that the muscle pair is mainly driving the stepping motion; CI $>1$ would mean that the muscle pair is resisting the motion; and CI $\sim 1$ means that the muscle pair either controls the motion or is not active.

**Statistical Analysis**

The motor errors, mean EMG, and co-contraction indices have a single value per stride (or step) for each subject. We used the SMART toolbox to report the errors and co-contraction values as continuous variables (van Leeuwen et al., 2019). The main advantages of using SMART over binning methods are that each subject contributes equally to the overall average and that the comparisons benefit from multiple-comparison correction and increased statistical power.

We used the Pingouin toolbox version 0.3.10 for our statistical analysis (Vallat, 2018). We compared motor errors, mean EMG, and co-contraction indices for each task at the end of pre, and the start and end of perturbed, catch, and post-perturbation blocks using repeated-measure analysis of variance (rANOVA). If the rANOVA was significant, we performed a priori post-hoc paired t-tests between the following pairs: 1) end of pre and end of the post to evaluate prolonged adaptation and sustained modifications 2) the start and end of perturbed strides to evaluate adaptation, 3) the start and end of catch strides to evaluate adaptation and anticipatory responses, 4) the start and end of the post to evaluate de-adaptation. We compared pre, and post-perturbation mean EMG and co-contraction indices between young and older adults using mixed-design ANOVA with age as
Figure 5.2: Schematic of motor errors and the motor error behavior for the left extension-onset and left mid-extension task blocks for young and older adults. For the far-left column, the vertical shaded light green rectangles indicate the period of the perturbations while the gray shaded box highlights the recovery step. The thick lines in the time courses are the mean and the shaded areas are the confidence intervals. * p<0.05. Horizontal brackets indicate significant differences from start to end. Vertical brackets indicate significant differences between end of pre and end of post.

a. The stepping duration (temporal) error was the difference between the duration of each step and the two-second mark set by the pacing cue (gray line). Young and older adults could maintain their temporal errors <100ms during the perturbed strides.

b. The maximum position (spatial) error was the maximum difference between each stride’s profile and the average baseline (pre) stepping profile. Spatial errors for young adults for the perturbed and catch strides did not converge by the end of the perturbation period whereas there was convergence for the older adults. Young and older adults sustained the increased spatial errors after the perturbations were removed during post (8-10 minutes) for both perturbation types.

The between-subject factor with two levels, young and old, and time as the within-subject factor, also with two levels, pre and post. We used Pingouin’s pairwise-ttest function value to 1 to determine if each muscle pair were driving (CI < 1), resisting (CI > 1), or controlling (CI ~1) the motion. With the automatic correction to perform the post-hoc tests if the mixed-design ANOVA was significant. We also compared the co-contraction index (CI) value to 1 to determine if each muscle pair were driving (CI < 1), resisting (CI > 1), or controlling (CI ~1) the motion. We used SMART’s
one-sample statistical test to determine if CI was different from one. The significance level of all statistical analyses was set to 0.05.

Results

Temporal error

Young and older adults did not reduce their temporal errors as they gained more experience with the perturbations, indicating a lack of error-based adaptation, but they did de-adapt after the perturbations were removed (Figure 5.2a). Both young and older adults had 50ms temporal errors during perturbed strides, while the temporal errors during catch strides were 150ms (Figure 5.2a). The rANOVA was significant for temporal errors in each task (young adults $F_{(6,96)} > 40, p < 0.0005$, older adults $F_{(6,96)} > 20, p < 0.0005$). However, the post-hoc a priori tests only indicated significant and meaningful differences in the left mid-extension temporal errors at start and end of catch strides for both young and older adults (young adults $p = 0.047$, older adults $p = 0.025$). Both young and older adults reduced their temporal errors to baseline levels after the perturbations were removed, indicating de-adaptation. The right-side showed also similar temporal errors, with the exception of young adults increasing their temporal errors form start to end of both perturbed and catch strides ($F_{(6,96)} = 144, p < 0.0005$, post hoc $p < 0.01$) (Supplementary Figure 5.6a).

Spatial error

Spatial errors of older adults during catch and perturbed strides converged by the end of the perturbed block whereas there was no convergence for young adults (Figure 5.2b). Spatial errors for young adults during the catch strides were $10^\circ$ for both perturbation tasks, which was less than $12^\circ$ for left extension-onset perturbed strides and $16^\circ$ for the left mid-extension perturbed strides. However,
the difference between the spatial errors of catch strides and of perturbed strides for older adults was not present in the left extension onset and diminished toward the end of the left mid-extension perturbations. The rANOVAs were significant for the spatial errors of every task (young adults $F_{(6,96)}>20$, $p<0.0005$, older adults $F_{(6,96)}>10$, $p<0.0005$). The post-hoc tests showed that the spatial errors for older adults were different from the start to the end of the catch strides for both left extension-onset and left mid-extension tasks (extension-onset $p = 0.023$, mid-extension $p = 0.057$). After removing the perturbations, spatial errors were always higher than pre levels for both young and older adults and did not return to pre levels (post hoc, young adults: extension-onset $p=0.018$, mid-extension $p=0.001$; older adults: extension-onset $p=0.008$, mid-extension $p=0.009$). For the right side tasks, spatial errors of catch strides for older adults did not increase from start to end (Supplementary Figure 5.6b). Nevertheless, spatial errors were also higher during the right-side task’s post blocks than the pre, but older adults (post hoc, young adults’ extension-onset $p=0.001$, mid-extension $p<0.0005$, older adults’ extension-onset $p=0.005$, mid-extension $p=0.002$).

*Muscle activity and co-contraction for the extension-onset perturbation tasks*

Young and older adults used their left (ipsilateral) rectus femoris and right (contralateral) anterior deltoid to drive the stepper during the left extension-onset perturbed steps (Figure 5.3). The agonist muscles (e.g., left rectus femoris and right anterior deltoid) were active mainly during the extension-onset, and the antagonist muscles (e.g., left semitendinosus and right rectus femoris) became active toward the end of the step, likely to slow down the stepping motion and assist with a smooth transition between steps (Figure 5.3, EMG profile). Of the six agonist muscles that young and older adults could use to drive the stepper, the left rectus femoris and the right anterior deltoid showed greater muscle activity during perturbed strides than catch strides (Figure 5.3, EMG mean). The mean left posterior deltoid antagonist activity increased during the recovery steps for young adults and did not return to the pre-perturbation levels after removing the perturbations (rANOVA
$F_{(6,96)}=2.4$, $p=0.01$, post-hoc perturbed stride $p=0.06$, pre-post $p=0.03)$. Older adults, however, showed greater mean EMG at the end of the post block compared to pre block for the left anterior deltoid, right anterior deltoid, and right soleus (rANOVA $F_{(6,60)}'>3.0$, $p's <0.01$, post hoc $p's <0.05$). Similar to the left side, right rectus femoris were active during the perturbed strides (Supplementary Figure 5.7). While most muscles had slightly increased their mean EMG, the right anterior deltoid mean EMG decreased for young adults during both perturbed and recovery steps and for older adults during the recovery step (rANOVA $F_{(6,60)}'>3.5$, $p's <0.01$, post hoc $p's <0.05$).

During the perturbed steps of the left extension-onset task, young adults only had CI ~1 for their right tibialis anterior-soleus while CIs for the other muscle pairs were <1 (SMART $p's<0.05$) (Figure 5.3, co-contraction index). During the recovery steps, CIs for left anterior deltoid-posterior deltoid and right anterior deltoid-posterior deltoid were ~1. Older adults, however, had CIs ~1 for all muscle pairs except for the left and right rectus femoris-semimembranosus during the perturbed step and left tibialis anterior-soleus during the recovery steps (SMART $p's<0.05$). Older adults reduced their co-contraction during the perturbation block for the left rectus femoris-soleus while using their right anterior deltoid-right posterior deltoid pair to resist the motion during the recovery steps (SMART $p's<0.05$). For the right extension-onset task, older adults also demonstrated similar CI reduction for the left anterior deltoid-posterior deltoid and briefly for the right rectus femoris-semimembranosus and the left and right tibialis anterior-soleus (Supplementary Figure 5.7).
Figure 5.3: EMG profiles (i.e. linear envelopes), time course of the EMG means for each block, and the time course co-contraction indices for the left extension-onset task for young (black headers) and older (gray headers) adults. The EMG profile is the grand average across the strides and subjects within the age-group. EMG mean and co-contraction is quantified per each perturbed and recovery step for the duration of the task. In the time courses, the thick lines are the mean and the shaded regions are the confidence intervals. * indicates significant post-hoc test. Colored rectangles at the bottom of the co-contraction plots indicate difference from 1 using SMART.
Muscle activity and co-contraction for the mid-extension perturbation tasks

Young adults used a greater number of agonists during left mid-extension than older adults (Figure 5.4). Young adults used the left (ipsilateral) posterior deltoid, left rectus femoris, left soleus, right (contralateral) semitendinosus, and right tibialis anterior to drive the stepper during the perturbed step (Figure 5.4, EMG profile). The antagonist muscle activity was dominant toward the end of the perturbed step, similar to the left extension-onset task. The left mid-extension perturbations resulted in increased left soleus activity for young adults during perturbed and recovery steps even after the perturbation was removed \((r\text{ANOVA} F'_{(6,96)}>2.3, p's<0.03, \text{post hoc} p's<0.05)\). Older adults used their left posterior deltoid, left rectus femoris, left soleus as agonists to overcome the left mid-extension perturbations (Figure 5.4, EMG profile). During the left mid-extension perturbation task, right and left rectus femoris and left posterior deltoid EMG activity significantly increased for older adults irrespective of their role in the movement \((r\text{ANOVA} F_{(6,54)}>4.3, p's<0.001, \text{post hoc} p's<0.05)\) (Figure 5.4, EMG mean). Young and older adults used largely the same muscles to overcome the right mid-extension perturbations, namely right (ipsilateral) posterior deltoid, left rectus femoris, right (contralateral) anterior deltoid. Older adults also continuously used their right soleus to drive the stepper which resulted increase in mean EMG for the perturbed and recovery steps during perturbed stride and form pre to post \((r\text{ANOVA} F_{(6,60)}>2.3, p's<0.04, \text{post hoc} p's<0.05)\) (Supplementary Figure 5.8).

The older adults had a significant CI increase during the perturbed step for the right anterior deltoid-posterior deltoid pair due to the increased antagonist right posterior deltoid became activity right after the perturbation (Figure 5.4, EMG profile; CI r\text{ANOVA} F_{(6,54)}=4.0, p=0.002, \text{post hoc} p's<0.01). Like the left extension-onset tasks, the Cis for muscle pair in young adults were <1 except for the anterior deltoid-posterior deltoids during the perturbation block (SMART p's<0.05). Older adults significantly reduced their CI during the perturbation block in the left anterior deltoid-posterior
deltoid, left rectus femoris-semitendinosus, and the right tibialis anterior-soleus to help drive the perturbed step (SMART p’s<0.05). For the right mid-extension task, older adults demonstrated a similar CI reduction for the right and left anterior deltoid-posterior deltoid, briefly for the left rectus femoris-semitendinosus, and the left and right tibialis anterior-soleus (Supplementary Figure 5.8).

Muscle activity and co-contraction differences pre to post for the extension-onset tasks

Overall, more muscles had significantly different amplitudes from pre to post in the left extension-onset task than the left mid-extension perturbations (Figure 5.5). The left posterior deltoid showed the most significant differences between young and older adults and the pre and post blocks during the left extension-onset task (mixed-design ANOVA F(1,26)’s>5, P’s <0.03, post-hoc, pre-post p’s<0.03). The mean EMG increased and was sustained from pre to post for the left posterior deltoid for both perturbed and recovery steps, and the mean EMG for young and older adults was also different in the perturbed (left) step (mixed-design ANOVA F(1,26)’s>4, p’s <0.03, post-hoc, pre-post p’s <0.01, age perturbed step p=0.02). These differences, however, only resulted in a significant CI increase from pre to post for the recovery steps in the left anterior deltoid-posterior deltoid in the left extension-onset task (Figure 5.5, co-contraction index). During the left extension-onset task, the left and right rectus femoris, semitendinosus, and soleus showed significantly greater mean EMG during the post block compared to pre block only when they were acting as antagonists (mixed-design ANOVA F(1,26)’s>4, P’s <0.03, post-hoc, pre-post p’s<0.05). As a result, post-perturbation CI was greater than pre CI for just the right rectus femoris-semitendinosus and right tibialis anterior-soleus pairs during the perturbed (left) step (mixed-design ANOVA F(1,24)’s>4, P’s <0.04, post-hoc, pre-post p’s,<0.04). Older adults had significantly higher CI than young adults in pre and post blocks for the right rectus femoris-semitendinosus pair in the left extension-onset task (mixed-design ANOVA F(1,26)=7.3, P=0.01, post hoc, age effect p=0.01, age x pre p=0.06, age x post p=0.02) (Figure 5.5, co-contraction index).
Figure 5.4: EMG profiles (i.e., linear envelopes), time course of the EMG means for each block, and the time course co-contraction indices for the left mid-extension task for young (black headers) and older (gray headers) adults. The EMG profile is the grand average across the strides and subjects within the age-group. EMG mean and co-contraction is quantified per each perturbed and recovery step for the duration of the task. In the time courses, the thick lines are the mean and the shaded regions are the confidence intervals. * indicates significant post-hoc test. Colored rectangles at the bottom of the co-contraction plots indicate difference from 1 using SMART.
Figure 5.5: Comparison of mean EMG and co-contraction between pre and post for the left-side tasks. * indicates post hoc significant test. Boxes expand between the first and third quartile. Whiskers are minimum and maximum after rejecting the outliers. Outlier rejection is only for representation and was not included in the statistical analysis.
During the left mid-extension task, the left and right rectus femoris mean EMG were greater during the post block than the pre block, resulting in increased CI from pre to post for right rectus femoris-semitendinosus pair (mixed-design ANOVA $F_{(1,24)}$’s $>4$, $p$’s $<0.04$, post hoc, pre-post $p$’s $<0.04$). The right posterior deltoid had greater antagonist activity during the post perturbation block for both young and older adults during the left mid-extension task, resulting in increased CI from pre to post for the right anterior deltoid-right posterior deltoid pair (mixed-design ANOVA $F_{(1,24)} =1.1$, $p =0.001$, post hoc, pre-post $p=0.002$). For right-side tasks, there was a main effect of time (pre-post) on the EMG amplitude and co-contraction index. Additionally, age also affected the EMG mean of several muscles, including left soleus and left semitendinosus, and the CIs of several muscle-pair, including right rectus femoris-semitendinosus (Supplementary Figure 5.9).

Discussion

We quantified and compared the motor error behavior, muscle activity responses, and co-contraction indices of young and older adults responding to perturbations during a seated locomotor task. Like young adults, older adults retained prolonged motor modifications after the perturbations were removed, suggesting that use-dependent learning also occurred for older adults. Unlike young adults, the spatial errors in catch and perturbed strides converged by the end of the perturbation block for older adults. Young adults used more muscle pairs than older adults to drive the stepper during the perturbed steps and the recovery steps. The co-contraction of older adults generally increased as a result of the perturbations. However, older adults could also effectively reduce their CI for select muscles and change the role of the muscle pairs from controlling to driving based on the task, such as with the left rectus femoris-semitendinosus during both left extension-onset and mid-extension perturbations. Despite the different task mechanics, the results demonstrate that perturbing the stepping patterns of older adults could promote a reduction of co-contraction for
select muscle-pairs based on the perturbation timing.

Use-dependent learning was clearly evident during these perturbation tasks in young and older adults, based on not just motor error behavior but also muscle activity and co-contraction indices. For young and older adults, motor errors did not decrease during the perturbed block, indicating that error-based adaptation did not occur. Instead of decreasing, the spatial errors were prolonged during the perturbed block and sustained through the post block in both young and older adults, which is indicative of use-dependent learning. This suggests that regardless of age, subjects perceived that following the pacing cue was their main goal in the perturbed stepping tasks and that modifying the stepping profile did not hinder achieving the task goal, which led to the retention of the modified stepping profile [21]. In general for young and older adults, muscle activity was higher during the post block after the perturbations were removed compared to the pre block, which provides more evidence of use-dependent learning. Further, the co-contraction indices did not decrease as subjects gained more experience with our perturbations or from pre to post. In typical error-based adaptation studies, co-contraction often decreases with adaptation [1, 38]. Taken altogether, motor errors, muscle activity, and co-contraction indices clearly indicate that use-dependent learning occurred as subjects responded to perturbations applied on a stride-by-stride basis during recumbent stepping.

Older adults used fewer muscle pairs to drive the stepper compared to young adults. Recumbent stepping is a mechanically redundant task. As such, subjects can drive the stepper with just one shoulder, elbow, hip or knee flexor-extensor muscle pair such as the left rectus femoris-left semitendinous pair or with a functional agonist-antagonist such as the left and right rectus femoris muscles that would oppose each other during stepping. In this task, the co-contraction index (CI) could delineate whether a muscle pair was driving (CI < 1), controlling (CI > 1), or refining (CI ~1) the stepping motion. Based on CI < 1, older adults had 3 out of 6 muscle pairs driving the stepping motion compared to the 5 out of 6 muscle pairs for young adults, indicating that older adults used fewer resources to drive the stepper. This aligns with previous studies of perturbed
walking and perturbed balance that showed that older adults used fewer muscle synergies to respond to the perturbations than young adults [39, 40]. Overall, by increasing co-contraction to potentially increase limb stiffness, older adults seemed to be able to resist and reject the perturbations such that the older adults had similar, if not smaller, motor errors during perturbed stepping compared to young adults. The only instance of a muscle pair significantly resisting the stepping motion was during the left extension-onset perturbations for the right anterior deltoid-posterior deltoid of just the older adults.

The reduction of co-contraction in select muscle pairs during perturbations is a novel aspect of this seated locomotor task. Interestingly, perturbation could decrease CI for select muscle pairs across both right- and left-side tasks (Figures 5.3,5.4 and supplementary Figure 5.7, 5.8). The CI decrease is in contrast to previous walking and balance studies which reported increased co-contraction with the presence of perturbations [13, 16, 17]. The perturbation timing resulted in different muscle pairs with reduced CIs for each task. This can be especially beneficial for rehabilitation, where using different perturbations during recumbent stepping would engage specific muscle-pairs to drive the locomotion. Recent studies have indicated that seated locomotor exercise significantly improves walking performance during rehabilitation [41, 42]. Transferability of perturbed seated exercises to walking improvement requires further research.

Limitations of this study include not including the handles and force data and attributing the perturbations to the extending leg. The recumbent stepper is equipped with load cells for pedals and handles. However, we decided not to use the force and moment data for this study because the inertia of the device would contaminate the force data, especially during the perturbations. While we asked subjects to use both arms and feet to drive the stepper, we attributed the perturbations to the extending leg. A previous study and our preliminary tests (not reported here) showed that the lower-limb extension contributes the most to compensate for increased stepping resistance [43].
Perturbed recumbent stepping generally increased co-contraction for both young and older adults from pre to post perturbation block and from the start to the end during the perturbation block. However, based on the timing of the perturbations, older adults reduced co-contraction of select muscle pairs to help drive the stepper. This novel effect and lack of fall risk provide the potential to use perturbations in seated locomotor tasks to help older adults reduce co-contraction and improve their motor response at the same time. Still, the efficacy of perturbed sitting tasks on walking performance and the effect of different perturbation intensity and duration require further research.

References


Supplementary figures

Figure 5.6: Motor errors for the right-side tasks
Figure 5.7: EMG profiles (i.e., linear envelopes), time course of the EMG means for each block, and the time course co-contraction indices for the right extension-onset task.
Figure 5.8: EMG profiles (i.e., linear envelopes), time course of the EMG means for each block, and the time course co-contraction indices for the right mid-extension task.
Figure 5.9: Comparison of mean EMG and co-contraction between pre and post for the right-side tasks.
CHAPTER 6: GROUP-LEVEL CORTICOMUSCULAR CONNECTIVITY DURING SEATED LOCOMOTOR PERTURBATIONS FOR YOUNG AND OLDER ADULTS

Introduction

Motor and locomotor perturbations elicit cortical and muscular activity, potentially reducing motor errors. Several cortical and subcortical brain areas are known to engage in monitoring motor errors. The anterior cingulate cortex in the mid-prefrontal brain region is likely to detect motor errors with sensory inputs and the expected goal [1]. The supplementary motor area integrates neural pathways from the motor cortex and anterior cingulate and likely acts as a sensory and executive hub for processing errors [2]. The posterior parietal cortex likely processes the differences between the visual and sensorimotor inputs [3]. Nevertheless, the activities of these brain areas are not essentially exclusive. For example, the left posterior parietal cortex has a critical role in joint impedance control, while the right prefrontal cortex controls the limb speed during perturbed upper limb motor tasks [4]. Perturbations tend to increase muscular activation and agonist/antagonist co-contraction [5, 6]. Not surprisingly, the cortico-cortical, intra-muscular, and corticomuscular coherence also increases with perturbations indicting increased cortical control over muscular activation [7–9].

The age-related differences in cortical and muscular activity are in different directions. Older adults implement more co-contraction in response to motor perturbations and would likely reduce their contraction less than young adults as they gain more experience with the perturbations [10]. However, locomotor perturbation studies also indicated that older adults use simpler muscular control through fewer muscle synergies or fewer driving muscles to maintain similar motor performance as young adults [11, 12]. Many studies have indicated “overactivation” of older adults’ brain regions compared
to young adults as a compensatory mechanism for similar motor performance [13, 14]. Such overactivation would increase connectivity at both cortical and muscular levels when facing motor challenges [15–17]. In contrast, a couple of studies on standing and locomotor tasks using channel EEG and EMG signals indicated weaker connectivity in older adults [18–20]. The different methods for connectivity computation, lack of source-resolved EEG, and inter-subject variability may contribute to the different outcomes observed in the connectivity studies [21, 22].

Nevertheless, the decrease in cortico-muscular connectivity due to neurological impairments is well-documented [23–25], and rehabilitation can improve the connectivity levels or use connectivity as a metric for motor improvement [26,27]. We previously introduced perturbed recumbent stepping as a locomotor task that can elicit similar brain dynamics as perturbed walking, especially in the anterior cingulate cortex and supplementary motor areas [28]. We also showed that older adults recruited fewer muscle-pairs than young adults to perform perturbed stepping, despite having similar motor errors as young adults. The main advantage of recumbent stepping and other seated exercises is that subjects would not require to maintain their balance while performing the exercise, which is especially beneficial for accessible walking rehabilitation. Determining the connectivity markers of perturbed stepping will set the baselines for using this exercise as a rehabilitation approach to improve corticomuscular connectivity.

The purpose of this study was to quantify the corticomuscular connectivity patterns of young and older adults in response to brief perturbations during recumbent stepping. We also aimed to determine whether the timing of the perturbations modulated corticomuscular connectivity. We hypothesized that the corticomuscular connectivity would increase around the perturbation timing, especially between the motor cortex and the driving muscles, namely, posterior deltoids and rectus femoris. This hypothesis was based on our previous EEG and EMG analyses showing increased electrocortical and muscular activation around the stepping perturbation [28]. We also hypothesized that the anterior cingulate cortex would have increased direct causal connectivity from a subset of
muscles around the perturbation timing, which would act as sensory feedback for error monitoring. In this study, we use connectivity and causal relation or effect interchangeably.

Methods

Seventeen adults (11 females, age 25 4.9 years) and 11 older adults (4 females, age 68 3.6 years) participated in the study. Subjects were all right-handed as they would pick an object from the floor, and they reported no prior neurological or physical problems for two years prior to their test. The Institutional Review Board of the University of Central Florida approved the study, and subjects gave their written informed consent before starting the experiment.

Hardware

We used a recumbent stepper integrated with a servomotor [29] to introduce brief perturbations in the form of added resistance during stepping. The stepper was mechanically coupled in a way that the contralateral arm and leg would extend together. We used the servomotor to briefly increase stepping resistance for 200 ms as the subject’s left or right leg was at the extension-onset or mid-extension. The increased stepping resistance was a torque-controlled algorithm that would require 3x the normal torque to drive the stepper at 60 steps per minute. We used the servomotor’s position sensor to record the stepper’s kinematics at 100Hz.

We used twelve wireless electromyography, EMG, sensors (Trigno, Delsys, Natick, MA) to collect muscular activity at ~1.1 kHz. We attached the EMG sensors on the tibialis anterior, soleus, rectus femoris, semitendinosus, anterior deltoid, and posterior deltoid on both left and right sides (Hermens et al. 1999). Here, we only analyzed the left-side muscles because our preliminary results indicated that including all muscles would decrease the connectivity accuracy. We used
high-density electroencephalography, EEG, system (ActiveTwo, 128 electrodes, BioSemi B.V.,
Amsterdam, the Netherlands) to collect brain activities at 512 Hz. We placed the EEG cap and
electrodes according to the BioSemi manual and used a 3D scanner (Structure I, Occipital Inc., CO)
to record the electrode’s 3D locations [30]. Data streams of the different systems were synchronized
using a trigger signal from the stepper controller to the EMG and EEG systems.

Protocol

Data collection consisted of four 10-minutes stepping tasks; each task only included one perturbation
type at the mid-extension or extension-onset of the left or right legs. Each task started and ended with
two-minutes pre and post unperturbed stepping blocks and included a six-minute-long perturbation
block, in which subjects would experience perturbation at each stride. There was also a random
one-in-five catch stride in the perturbation block, which did not include perturbations. Before
starting each task, we asked subjects to 1) follow the visual pacing cue presented in front of them
at 60 steps per minute, 2) step smoothly as they were walking, and 3) use both arms and legs
to drive the stepper. A complete explanation of the protocol has been presented previously [28].
We determined the stepping events using the kinematic information from the stepper servomotor.
Each stride was from the start of the perturbed-leg extension to the next perturbed-leg extension.
Perturbation events were the start of the increased stepping resistance for each stride.

EEG processing

EEG and EMG data were processed and analyzed in MATLAB environment (R2018b, MathWorks
Inc., Natick, MA) with a custom pipeline based on EEGLAB version 2019.0 [31] (Figure 6.1). The
EEG pre-processing pipeline is largely similar to our previous study on young adults’ perturbed
stepping [28]. In summary, we used a high-pass filter at 1 Hz and a 60 Hz line-noise filter to clean the
EEG data before applying the template correlation rejection method to identify the channels that had a high correlation to the stepping events [32]. We then used step-wise channel and frame rejection to create 32 datasets for each subject with a range of conservative to lenient noise rejection. The signal noise rejection metrics were range, variance, kurtosis, and correlation to other channels [33]. The frame noise rejection metric was the EEG inter-quartile variability to the overall median of the signal during each task. We used adaptive mixture independent component analysis (AMICA) to quantify the independent components (ICs) for each of the 32 datasets, estimated the IC locations in the brain using DIPTFIT, and used a multivariate classifier (ICLabel) to estimate the number of "brain" ICs for each dataset [34, 35]. We ultimately the dataset with the most "brain" ICs at the representative dataset for each subject.

We clustered ICs from young and older adults separately based on the IC location, power spectrum, and scalp map. We used the 3 to 25 Hz range for power spectrum and the Laplacian of the scalp map to perform the clustering [36]. We used optimal k-means clustering to select the number of cortical clusters for young and older adults separately and focused on the clusters that included >70% of the subjects. If the ICs in a cluster were mostly in a Brodmann area, we would assign that
cluster with that Brodmann area [30, 37]. If a cluster spanned across multiple Brodmann areas, we would attribute the cluster with the region that would encompass the cluster’s ICs.

Finally, we computed the temporal profile of the spectral power of each cluster, commonly known as the event-related spectral perturbations (ERSP) [38]. We epoched the EEG data from -400 ms to +400 ms of the perturbation events. We baseline normalized the spectral power using the average spectral power during pre and computed the statistically significant event-related synchronization and desynchronization for the ERSPs using EEGLAB permutation functions with alpha equal 0.05 [39]. ERPS images only show significant spectral fluctuations.

**Connectivity processing**

We used the cortical source clusters and EMG from twelve upper and lower limb muscles to compute connectivity metrics mainly based on Peterson and Ferris’s group-level connectivity analysis of balance-beam walking and standing perturbations [2] (Figure 6.1). We selected each subject’s ICs that contributed to the cortical clusters and added the EMG signals after high-pass filtering at 1 Hz and rectifying [40]. We epoched the integrated IC and EMG data from -2 sec to +2 sec of the perturbation events to provide the required data for estimating the multivariate autoregressive models for each subject. We used the Viera-Morf algorithm with 1.3-sec window size and 0.01-sec window step-size to fit a multivariate autoregressive model with an order equal to 32 to each subject’s integrated dataset [41, 42]. We used the 1.3-sec window size to improve the model fit and the 0.01-sec window step-size to provide 10 ms temporal resolution for the connectivity analysis. The model order of 32 provides a spectral resolution of 4 Hz for the connectivity analysis and lets the autoregressive model take more of the past data into the model. The higher model orders, up to 40, have not shown significant noise increases inherent to estimating a greater number of variables in a similar study [2].
We used the direct Directed transfer function (dDTF) to estimate the direction connectivity between cortical ICs and EMG signals. The main advantages of dDTF are the casual directionality and the separation of direct causal effects from indirect causal effects [2,42–44]. Casual directionality provides the source and target of the "information flow" instead of the mere coherent activation of two sources [44]. Exclusion of the hierarchy effect where a source upstream has a causal effect on all downstream targets is another advantage of dDTF because only the direct causal influences are taken into account [43]. A phantom-head study with motion artifacts also had shown that dDTF could effectively recover the casual relations between signal sources [45]. We finally averaged the cortical clusters and EMG source connectivity results across subjects, baseline normalized to the average connectivity from -400 ms to -100 ms seconds and showed the significant increases and decreases in connectivity similar to the ERSP plots from -400 ms to +400 ms of the perturbation event.

We also compared the baseline corticomuscular connectivity between young and older adults to determine the pre-perturbation coherence across tasks. We averaged the brain to muscle and muscle to brain connectivity values across cortical clusters and muscles, and also across the frequency spectrum for the left-side tasks to quantify the overall directional connectivity. We compared the baseline connectivity using the Mixed-design ANOVA with they task type (left extension-onset or left mid-extension) and age (young or older adults) as the independent variables and connectivity direction (brain to muscle or muscle to brain) as the dependent variable. The significant level for all statistical tests was set at 0.05.

Results

Young adults’ EEG resulted in more individual “brain” ICs and more group-level clusters than older adults (Figure 6.2). The median “brain” component for young adults was 22 per subject, which
Figure 6.2: The location of the group cortical clusters for young and older adults. Older adults did not have an anterior cingulate cortex or differentiated motor cortex cluster. Young adults had strong theta-band (3-8Hz) synchronization locked to the perturbations in the

was significantly greater than the 10 per subject median for older adults (Wilcoxon rank-sum test, p< 0.0005). Young adults had five clusters with ICs from> 70% of the subjects. The clusters were at the anterior cingulate cortex, right supplementary motor area, left supplementary motor area, right posterior parietal cortex, and left posterior parietal cortex. Older adults had only three clusters with ICs from > 70% of the subjects. Since the ICs within the clusters were more spread to assign a Broadmann area to the clusters, only the general cortical area was attributed to the older adult clusters. The clusters were at the motor cortex, left posterior parietal cortex, and right posterior parietal cortex.
Figure 6.3: The spectral fluctuations (ERSP) image around the perturbation even (P) for young and older adults during the left extension-onset. Young adults’ Anterior cingulate cortex, supplementary motor areas, and left posterior parietal cortex had significant theta-band (3-8 Hz) synchronization after the perturbation event. Older adults had minor theta and alpha (3-8 Hz) synchronization in their motor cortex and left posterior parietal cortex.

anterior cingulate and supplementary motor areas for both left extension-onset. However, older adults only showed slight theta synchronization after the perturbations in the motor cortex (Figure 6.3a). The left posterior parietal cortex had increased theta and alpha synchronization for both young and older adults after the perturbation event. However, the right posterior parietal cortex did not show significant spectral fluctuations locked to the perturbation in either young and older adults (Figure 6.3a,b). Both young and older adults also demonstrated theta-alpha desynchronization before the perturbation event in the left posterior parietal cortex.

Mid-extension perturbations also elicited theta-band synchronization in the young adults’ anterior cingulate and right and left supplementary motor areas (Figure 6.4a). Older adults, however, only had slight theta-band motor cortex synchronization toward the end of the 400 ms window (Figure 6.4b). The left posterior parietal cluster had a beta-band (13-35 Hz) synchronization before perturbation which was followed by a beta desynchronization after the perturbations. Older adults
Figure 6.4: The spectral fluctuations (ERSP) image around the perturbation even (P) for young and older adults during the left mid-extension. Young adults’ Anterior cingulate cortex and supplementary motor areas had significant theta-band (3-8 Hz) synchronization after the perturbation event. Older adults had minor theta and alpha (3-8 Hz) synchronization in their motor cortex and right posterior parietal cortex at +400ms.

did not show significant posterior parietal spectral fluctuations before or after the perturbation event.

The baseline connectivity analysis revealed significant directional and age effects on the causal realtion between the brain and muscles (Figure 6.5). Older adults had greater baseline connectivity in both brain to muscle and muscle to brain directions (mixed ANOVA, $F_{(1,944)}=14.1$, $p<0.0005$). Also, muscle to brain connectivity was weaker than the brain to muscle connectivity (mixed ANOVA, $F_{(1,944)}=36.0$, $p<0.0005$). The task type, however did not have a significant effect on the directional connectivity (mixed ANOVA, $F_{(1,944)}=2.9$, $p<0.087$).

Brain to muscle connectivity analysis showed selective anterior cingulate causal muscle relations in young adults and the left parietal cortex causal relation to all muscles for older adults (Figure 6.6). The anterior cingulate elicited left soleus activity after the perturbations in the left extension-onset perturbations but mostly elicited the left posterior deltoid for left mid-extension perturbations.
Figure 6.5: The baseline connectivity prior to the perturbation onset. Older adults had significantly greater baseline connectivity than young adults (shown by asterik). Also, brain to muscle connectivity was significantly greater than the muscle to brain connectivity (shown by hash).

(Figure 6.6a,b). The right supplementary motor area had a consistent causal connection to all left-side muscles after the perturbation event for young adults. Compared to the left mid-extension, the left supplementary motor area and right posterior parietal cortex seem to elicit more activity in the anterior and posterior deltoid during left extension-onset perturbations. The motor cortex and the left posterior parietal cortex of the older adults elicited muscular activity to all left-side muscles (Figure 6.6c,d). The older adults’ right posterior parietal cortex seemed to have stronger causal connections to the soleus and rectus femoris for left extension onset and to the posterior deltoid for the left mid-extension perturbations.

The lower-limb muscles had an increasing causal relationship to the anterior cingulate activity in young adults, and most older adults’ muscles had an increasing causal relation to the motor cortex during and around the perturbations (Figure 6.7). The left-side tibialis anterior, soleus, and rectus femoris had increased connectivity to the anterior cingulate cortex around the left extension-onset perturbations. On the other hand, only the left rectus femoris had strong increasing connectivity to the anterior cingulate after left mid-extension perturbations. The left and right supplementary motor
areas had increased connectivity from the anterior deltoid in the left-extension-onset, but the left rectus femoris was the common, increasing connectivity for the left and right supplementary motor areas for the mid-extension perturbations. Interestingly, the anterior deltoid connectivity to the right supplementary motor area decreased after the left mid-extension perturbations. All muscles had increased causal relation to the older adults’ motor cortex during and after the left mid-extension perturbations (Figure 6.7c). However, the mid-extension perturbations did not increase left tibialis anterior and posterior deltoid connectivity to the motor cortex (Figure 6.7d). All the left-side muscles also had increased connectivity to the left or right posterior parietal cortex for older adults.

Discussion

We quantified the electrocortical spectral dynamics and corticomuscular connectivity around perturbations during recumbent stepping for young and older adults. Young adults had concentrated electrocortical sources in the anterior cingulate, left and right supplementary motor area, and left and right posterior parietal cortex. The anterior cingulate and supplementary motor area clusters consistently showed strong theta-band synchronization after the perturbation event. Older adults, however, had fewer electrocortical sources, especially in the anterior cingulate cortex and supplementary motor area region. Also, older adults only showed slight significant synchronization after the perturbations compared to unperturbed stepping, suggesting that the older adults’ electrocortical activity was not significantly different from the unperturbed stepping. Directional causal connectivity results showed a subset of muscles were directly influenced by the anterior cingulate activity, and only a subset of muscles had a direct influence on the anterior cingulate activity. Interestingly, older adults increased their corticomuscular connectivity after the perturbation and had consistent left posterior parietal to muscle connectivity during both tasks. The results provide more evidence of significant corticomuscular connectivity despite the plateaued cortical activity for older adults.
Figure 6.6: Brain to muscle connectivity. Young adults’ Anterior cingulate cortex had strong causal relation with the left soleus in the left extension-onset and with posterior deltoid in the left mid-extension. Older adults had consistent causal relation from the motor cortex to most of the left-side muscles during both tasks. The left posterior parietal cortex had strong connectivity to older adults’ muscles in both tasks. ACC = Anterior cingulate cortex, r. & l. SMA = right and left supplementary motor area, r. & l. PPC = right and left posterior parietal cortex, TA = tibialis anterior, SO = soleus, RF = Rectus femoris, ST = semitendinosus, AD = anterior deltid, PD = posterior deltoid.
Older adults had fewer electrocortical ICs per subject and fewer cortical group clusters than young adults. Previous neuroimaging studies have suggested overactivation of the brain areas in older adults for performing a similar motor task as young adults [13]. Such overactivity results in several brain regions becoming active, likely with similar signal characteristics [14]. The activity of the overactive regions may not be temporally independent, resulting in shared ICs for different regions, which in turn, would be rejected by DIPFIT algorithm because one of the basic assumptions of the inverse solution is that the brain ICs are dipolar. Further, activity synchrony of different regions of the brain would create a global signal pattern that, with common average referencing, would be canceled from the signal during preprocessing [18, 46].

Young adults had selective information flow to and from the anterior cingulate, but older adults did not present anterior cingulate activity with the presence of the perturbations. The anterior cingulate cortex compares the expected motor goals with the sensory inputs [1, 47], so we hypothesized that part of the sensory input would come from a subset of muscles. We found that during extension-onset tibialis anterior, soleus, and rectus femoris elicit anterior cingulate activity, while during mid-extension perturbations, rectus femoris has the most significant connectivity to the anterior cingulate (Figure 6.7a,b). Interestingly, upper limb muscles (left anterior deltoid and posterior deltoid) did not contribute to the anterior cingulate activity. Older adults, however, lacked a group cluster located at the anterior cingulate cortex. Looking at our previous research on muscular co-contraction in the previous chapter, both young and older adults generally used their lower-limb muscles to drive the stepper during the perturbed step. So, the lack of anterior cingulate response in older adults might be beyond the lack of sensory input.

Older adults still maintained corticomuscular connectivity despite fewer cortical areas involved in the process. The left posterior partial connectivity to muscles seemed stronger for older adults. While older adults had only one overall cortical cluster at the motor cortex, almost all left-side muscles seem to have strong bi-directional connectivity with that cluster (Figure 6.6c,d and Figure 6.7c,d).
Figure 6.7: Muscle to brain connectivity. Young adults’ lower-limb muscles had causal relations to the anterior cingulate cortex. Older adults had consistent casual relation from the muscles to motor cortex. TA = tibialis anterior, SO = soleus, RF = Rectus femoris, ST = semitendinosus, AD = anterior deltoid, PD = posterior deltoid, ACC = Anterior cingulate cortex, r. & l. SMA = right and left supplementary motor area, r. & l. PPC = right and left posterior parietal cortex.

This strong connectivity is in line with the previous studies indicating increased corticomuscular connectivity for challenging motor tasks, especially from the motor cortex [16, 17]. Interestingly, the left posterior parietal cortex had a stronger causal relation to older adults’ muscles across both left mid-extension and extension-onset tasks than young adults. Studies with stroke patients had previously indicated the role that the left posterior parietal cortex has in joint impedance control.
during perturbed upper limb tasks [4, 48]. The more active left posterior partial connectivity and subsequently impedance control in older adults might explain why older adults used fewer muscle pairs to drive the stepper (i.e., older adults had fewer muscle-pairs with agonist-dominant response).

Limitations of this study include a low number of unperturbed strides and a lack of cross-frequency coherence analysis. Despite acceptable temporal and frequency resolution, dDTF apparently requires many trials, at least more than 100 trials, to yield to acceptable results based on previous locomotor studies [2, 49]. There were ~30 catch strides in our study and ~50-60 strides during either the pre or post blocks. These limited strides were insufficient for the dDTF connectivity analyses. Alternative connectivity methods such as wavelet analyses may be better suited to analyze lower numbers of trials. The majority of our young adults’ motor cortex responses were at the theta-band, while the muscular activity had high beta and gamma (20+Hz) frequency activities. This disparity in frequency response might skew dDTF and similar connectivity approaches, which by design look for a causal relation between signals in a similar frequency range [43]. Also, the 4Hz-frequency resolution determined by the dDTF results in just two data points for the theta-band (3-8 Hz) connectivity. Treating different frequency bands as separate signals and fitting multivariate autoregressive models with different orders might result in a stronger representation of theta-band responses in the connectivity analysis.

Overall, the stepping perturbations increased bi-directional corticomuscular connectivity in young and older adults. Despite finding fewer cortical clusters for the older adults, all muscles had a significant causal relation to the motor left posterior parietal cortices; the posterior parietal cortex is responsible for joint impedance control, suggesting that impedance control might be a stronger motor control paradigm for older adults. The results provide new evidence that modifying perturbation timing can change connectivity and enhance specific corticomuscular pathways. Such corticomuscular enhancements can be beneficial in providing closed-loop and personalized rehabilitation [50].
References


APPENDIX A: IRB DOCUMENTS
CLOSURE

December 4, 2020

Dear Helen Huang:

On 12/4/2020, the IRB reviewed the following protocol:

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<tr>
<th>Type of Review:</th>
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<td>Brain and muscle responses to stepping errors during rhythmic exercise</td>
</tr>
<tr>
<td>Investigator:</td>
<td>Helen Huang</td>
</tr>
<tr>
<td>IRB ID:</td>
<td>CR000000840</td>
</tr>
<tr>
<td>Funding:</td>
<td>Name: National Institute on Aging (NIA), Funding Source ID: R01AG054621</td>
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<td>Grant ID:</td>
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<tr>
<td>IND, IDE, or HDE:</td>
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The IRB acknowledges your request for closure of the protocol effective as of 12/4/2020. As part of this action:

- The protocol is permanently closed to enrollment.
- All subjects have completed all protocol-related interventions.
- Collection of private identifiable information is completed.
- Analysis of private identifiable information is completed.

If you have any questions, please contact the UCF IRB at 407-823-2901 or irb@ucf.edu. Please include your project title and IRB number in all correspondence with this office.

Sincerely,

Racine Jacques, Ph.D.
Designated Reviewer
November 27, 2019

Dear Helen Huang:

On 11/27/2019, the IRB reviewed the following submission:

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<td>Brain and muscle responses to stepping errors during rhythmic exercise</td>
</tr>
<tr>
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<td>Documents Reviewed:</td>
<td>r01-rs-ad.docx, Category: Recruitment Materials; r01-rs-bbs.pdf, Category: Test Instruments; r01-rs-champs.pdf, Category: Survey / Questionnaire; r01-rs-consent-c1.pdf, Category: Consent Form; r01-rs-email.docx, Category: Recruitment Materials; r01-rs-flyer.doc, Category: Recruitment Materials; r01-rs-msse.pdf, Category: Test Instruments; r01-rs-phonesurvey.doc, Category: Survey / Questionnaire; r01-rs-protocol-c1.doc, Category: IRB Protocol; r01-rs-sppb.doc, Category: Test Instruments; r01-rs-subjinfo.docx, Category: Survey / Questionnaire;</td>
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In conducting this protocol, you are required to follow the requirements listed in the Investigator Manual (HRP-103), which can be found by navigating to the IRB Library within the IRB system.

If you have any questions, please contact the UCF IRB at 407-823-2901 or irb@ucf.edu. Please include your project title and IRB number in all correspondence with this office.

Sincerely,

Adrienne Showman
Designated Reviewer
MODCR00000307: Modification and Continuing Review #1 for Study Brain and muscle response stepping errors during rhythmic exercise

Principal investigator: Helen Huang
Submission type: Modification and Continuing Review
Primary contact: IRB office: UCF IRB
IRB coordinator: Adrienne Showman
Letter: Correspondence_for_Pre-2018_Requirements

History
Contacts
CITI Training
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Principal Investigator

<table>
<thead>
<tr>
<th>Name</th>
<th>Financial Interest</th>
<th>E-mail</th>
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<tbody>
<tr>
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<td>no</td>
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</tbody>
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Study Team

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<tr>
<th>Name</th>
<th>Roles</th>
<th>Financial Interest</th>
<th>Involved in Consent</th>
<th>E-mail</th>
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<tbody>
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<td>Cesar Castano</td>
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<td>Research Assistant</td>
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</tbody>
</table>

Other Study Team Member Information

Guests Who Can View This Submission

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<th>Name</th>
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Hi Seyed,

I have reviewed your dissertation document as well as the previously approved study, SBE-17-12863. I can confirm that your research methods were part of this approved protocol and that you were listed as a study team member. The research done in your dissertation was covered under IRB protocol SBE-17-12863.

Please let me know if I can assist further.

Regards,

Gillian Bernal, CIP
IRB Manager
Office of Research and Commercialization
University of Central Florida
Gillian.Bernal@ucf.edu
IRB Main Phone: 407-823-2901

Human Subjects Research during Covid-19:
https://corona.research.ucf.edu/human-subjects-research/

Good evening,

I hope you are doing fine and this letter finds you well. At the request of UCF Dissertation Services and the confirmation of Dr. Helen Huang, the PI of the IRB protocol SBE-17-12863 (please see below), please kindly issue an approval letter that states “the research done in [my] dissertation was covered under that IRB protocol.” I am listed as a member of the study team for this protocol. My dissertation is accessible from OneDrive [from this link].

I appreciate your help with this matter.

Best,
Seyed Yahya Shirazi
Hi Seyed,

As the PI of the approved and now closed IRB protocol SBE-17-12863, “Brain and muscle responses to stepping errors during rhythmic exercise,” I have written this email to confirm that the experiments in your (Seyed Yahya Shirazi’s) PhD were conducted according to IRB protocol SBE-17-12863 for which you were a research assistant. Racine Jacques (Racine.Jacques@ucf.edu) was the IRB coordinator for the protocol.

Thanks,
Helen
APPENDIX B: COPYRIGHT PERMISSION LETTERS
We, Seyed Yahya Shirazi and Dr. Helen J. Huang, the copyright holders of the paper entitled *Differential theta-band signatures of the anterior cingulate and motor cortices during seated locomotor perturbations* grant the permission to reprint this paper in Seyed Yahya Shirazi’s dissertation entitled *Corticomuscular adaptation to mechanical perturbations in a seated locomotor task*.

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Paper’s full citation:


Seyed Yahya Shirazi

Dr. Helen J. Huang
Assistant Professor
Department of Mechanical and Aerospace Engineering
University of Central Florida
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