Evidence for Motor Cortex Dedifferentiation in Older Adults

Jessica A. Bernard¹ & Rachael D. Seidler¹.²

¹Department of Psychology, University of Michigan
1012 East Hall, 530 Church Street
Ann Arbor, MI 48109-1043 USA
²School of Kinesiology, University of Michigan
401 Washtenaw Avenue
Ann Arbor, MI 48109-2214 USA

Key Words: aging, transcranial magnetic stimulation, motor cortex

*Corresponding author:
Jessica A. Bernard
530 Church St.
Ann Arbor, MI 48109-1043
Tel: (734) 764-8186
Fax: (734) 936-1925
Email: jessbern@umich.edu
Abstract

Older adults (OA) show more diffuse brain activity than young adults (YA) during the performance of cognitive, motor, and perceptual tasks. It is unclear whether this over-activation reflects compensation or dedifferentiation. Typically, these investigations have not evaluated the organization of the resting brain, which can help to determine whether more diffuse representations reflect physiological or task-dependent effects. In the present study we used transcranial magnetic stimulation (TMS) to determine whether there are differences in motor cortex organization of both brain hemispheres in YA and OA. We measured resting motor threshold, motor evoked potential (MEP) latency and amplitude, and extent of first dorsal interosseous representations, in addition to a computerized measure of reaction time. There was no significant age difference in motor threshold, but we did find that OA had larger contralateral MEP amplitudes and a longer contralateral MEP latency. Furthermore, the spatial extent of motor representations in OA was larger. We found that larger dominant hemisphere motor representations in OA were associated with higher reaction times, suggesting dedifferentiation rather than compensation effects.
1. Introduction

Substantial evidence has accumulated over the last decade demonstrating that task-related brain activity is more diffuse and bilaterally symmetrical in older than young adults (YA; cf. Cabeza, 2002; Reuter-Lorenz and Lustig, 2005). While the bulk of this work has been targeted at understanding age differences in cognitive function, a burgeoning literature documents that similar over-activation is observed when participants perform motor tasks (Mattay et al., 2002; Naccarato et al., 2006; Ward & Frackowiak, 2003; Riecker et al., 2006; Ward, 2006; Calautti et al., 2001; Hutchinson et al., 2002; Talelli et al., 2008a; Heunickx et al., 2005; 2008; Langan et al., 2010). Debate lingers as to whether and how such over-activation is related to changes in brain structure, function, and biochemistry with age, as well as whether more diffuse activation patterns reflect compensatory processes or dedifferentiation of (i.e., less distinctive) representations (for a recent review see Seidler et al. 2010).

Increased activation in prefrontal regions during the performance of cognitive tasks has been shown to improve performance in older adults (OA) suggesting that, at least under some circumstances, additional activation is compensatory (Cabeza 2002; Carp et al., 2010). Dedifferentiation, as defined by excess (less distinct) brain recruitment in OA (Park et al., 2001) is not mutually exclusive from compensation, and is frequently reported in the literature on cognitive aging. In visual areas there is evidence for dedifferentiation such that OA have less specific neural representations for different categories of visual stimuli (Park et al., 2004; Carp et al., In Press). Furthermore, recent work by Carp and colleagues (2010) has shown decreased distinctiveness in visual regions of the brain during the performance of a working memory task. The visual cortex voxels activated for verbal and visuospatial working memory tasks were overlapping. In the same participants performing the same tasks, activity in prefrontal regions during maintenance and retrieval was indicative of compensatory processes. The pattern of findings for motor tasks is less consistent. While some work suggests that bilateral activation in the motor cortex is compensatory (Mattay et al., 2002) resulting in better performance, there is
also evidence demonstrating that this overactivation is associated with poorer motor task performance (Langan et al., 2010). Any excess recruitment in OA that results in behavioral deficits will be referred to here as dedifferentiation, whereas compensation will refer to any excess recruitment that results in performance benefits.

More symmetrical activation of the motor system is predominantly associated with increased engagement of the motor cortex ipsilateral to the moving hand in OA (Mattay et al., 2002; Ward & Frackowiak, 2003; Heuninckx et al., 2005, 2008). Given that OA have reduced interhemispheric inhibition between the motor cortices compared to YA (Talelli et al., 2008a, 2008b), greater activation in the ipsilateral motor cortex may be a result of release from inhibition by the contralateral motor cortex. Indeed, Talelli and colleagues (2008a, 2008b) have demonstrated that OA do not modulate interhemispheric inhibition when going from a state of rest to muscle contraction to the same extent that YA do. Moreover, the authors found that the degree of change in interhemispheric inhibition from rest to muscle contraction was associated with the extent of fMRI activation in the ipsilateral sensorimotor cortex, regardless of age (Talelli et al. 2008a).

Less lateralized brain processing has been shown to be associated with reductions in callosal cross sectional area in OA as well (Müller-Oehring et al., 2007). It is unclear, however, whether greater interhemispheric crosstalk and more diffuse motor recruitment reflect brain organizational features that are present at rest, or are instead only associated with task performance. For example, it has been shown that recruitment of ipsilateral motor cortex increases with task difficulty in YA (Seidler et al. 2004; Verstynen et al., 2005). Greater ipsilateral motor cortical activation in OA may in part be a reflection of heightened task difficulty for older versus young adults. Or, it may be due to a reduced ability to modulate inhibitory processes to suit varying task demands (Talelli et al., 2008a, 2008b; Fujiyama et al., 2009).

Thus, the primary purpose of the current study was to compare motor cortical representations of the first dorsal interosseous (FDI) finger muscle between YA and OA and to
investigate whether the extent of these representations influences motor performance. We applied transcranial magnetic stimulation (TMS) to the motor cortex when participants were at rest. We hypothesized that OA would exhibit a greater number of ipsilateral motor evoked potentials (MEPs) elicited by TMS, which are typically not observed in the resting state in healthy YA (although we have recently shown that they occur in some YA, particularly those with low handedness scores; Bernard et al., 2011). We also predicted that the spatial extent of motor representations in OA would be more extensive than that of YA. Additionally, we predicted that these more diffuse representations in OA would have a negative impact on motor performance resulting in higher reaction times, consistent with the pattern of dedifferentiation seen in the motor cortex with age that we have recently reported (Langan et al., 2010).

A secondary purpose of this study was to evaluate whether there are age differences in MEP amplitude and latency. It is important to delineate age differences in motor cortical excitability and motor map characteristics because reorganization (expansion and contraction) of such maps is routinely tracked during the time course of physical rehabilitation interventions in patients who have suffered from stroke, who have Parkinson's disease, or other neurological impairments that exhibit increasing frequency with age. Moreover, the reorganization capacity of the motor cortex has been shown to vary with the MEP amplitude that can be evoked prior to intervention (Pitcher & Miles, 2002). Previous work has shown that OA have reductions in motor cortical plasticity based on studies using paired-associative stimulation (Tecchio et al., 2008) and motor training (Sawaki et al., 2003; Rogasch et al., 2009). The literature is mixed, however, in terms of whether OA have smaller MEP amplitudes than YA. Some studies have reported smaller MEPs in OA (Fujiyama et al., 2009; Oliviero et al., 2006; Talelli et al., 2008b), while others have found no age differences in MEP amplitudes (Tecchio et al., 2008).

We also wished to evaluate whether motor threshold differs with age. Some studies have reported no difference (Fujiyama et al., 2009; Pitcher et al., 2003; Talelli et al., 2008b; Rogasch et al., 2009), while others have reported that OA have lower motor thresholds than YA.
(Tecchio et al., 2008) or higher motor thresholds (Rossini et al., 1992; Oliviero et al. 2006 report a nonsignificant trend in this direction). These conflicting results may be due to the broad age range of OA tested across these studies (55 years up to 94 years old). Interestingly, in their sample of participants aged 70 to 94 years old, Silbert et al. (2006) found that resting motor threshold decreased with age, and lower thresholds were associated with increased white matter hyperintensities and increased cerebrospinal fluid volume. A larger scalp to cortex distance was associated with a larger resting motor threshold as well (Silbert et al., 2006). It is therefore of interest to compare motor threshold between young and older participants of a more restricted age range.

In the current study we mapped M1 representations of the first dorsal interosseous muscle in YA and OA in both the dominant (left) and nondominant (right) hemisphere. We recorded MEPs bilaterally allowing us to quantify resting motor threshold, ipsilateral and contralateral MEP amplitudes and latencies, and the extent of contralateral and ipsilateral motor representations. We hypothesized that OA would have more diffuse motor representations than YA, and would have a greater propensity to exhibit ipsilateral representations. We also predicted that OA would have smaller MEP amplitudes and larger resting motor thresholds.

2. Methods

2.1 Participants

16 young adult (21 ± 1.83 years, 4 male) and 17 older adult (69.53 ± 4.05 years, 7 males) participants were recruited from the University of Michigan and the surrounding community and were paid for their participation. Participants responded to recruitment fliers placed around the community and on an online recruitment website. Additionally, we directly recruited individuals interested in participating in research through a database. A subset of the data from the YA was previously reported (Bernard et al., 2011). Upon enrollment in the study participants signed an IRB-approved informed consent form. Exclusion criteria included a history of neurological disease or damage, migraines, arthritis, head injury, or psychiatric
disorder. All participants were cognitively healthy as measured by the Mini-Mental State Exam (MMSE; Folstein et al., 1975; mean ± SD; YA= 29.69 ± .60, OA=28.76 ± 1.64) and the Mattis Dementia Rating Scale (Mattis, 1976; YA=142.87 ± 1.20, OA=142.23 ± 1.56). Participants were all strongly right-handed based on their laterality scores from the Tapping Circles portion of the Hand Dominance Test (Steingrüber, 1971), which we have previously shown are correlated with the number of ipsilateral MEPs in YA (Bernard et al, 2011).

2.2 Experimental Set-Up and Procedure:

Testing occurred on two separate days. During one session participants completed the Edinburgh Handedness Inventory (Oldfield, 1971) to provide a self-report measure of handedness. We administered the Tapping Circles and Tapping Squares from the Hand Dominance Test (Steingrüber, 1971) to provide a graphomotor measure of handedness and dexterity for each hand. Participants had 30 seconds to tap on small circles or squares using a marker with their left and right hands. The order of hands was counterbalanced across participants. The number of circles and squares tapped with each hand was counted and used to calculate a laterality index. Participants also performed the Purdue Pegboard Test (right hand, left hand, bilateral, and assembly tasks, 30 seconds per trial, 3 trials per task; Tiffin & Asher, 1948) and a grip strength assessment. For the Edinburgh Handedness Inventory, the Hand Dominance Test, and the unimanual conditions of the Purdue Pegboard, we calculated laterality indices in the following manner: (R-L)/(R+L), where R and L indicate right and left hand responses, respectively. Finally, we measured reaction time of the dominant (right) hand using a computer task implemented with E-Prime 1.1 software (Psychology Software Tools, Inc). Participants responded with their right hand as quickly as possible to white dots (subtending 0.61° of visual angle) presented on a black background in either the left or right visual field (center of circle subtending 6.02° of visual angle to the left or right of a central fixation cross). The fixation cross was presented for a variable foreperiod (500, 650, 800, or 1000 ms) to
prevent anticipatory responses. Left and right visual field presentations were pooled for the calculation of reaction time (RT) and any responses under 100 ms were eliminated.

On the second test day, participants underwent a TMS motor-mapping procedure. All participants were screened for any potential contraindications for TMS, following the guidelines of Rossi et al. (2009). Participants sat comfortably in a chair with their head resting in a chin rest and their hands relaxed. A tight-fitting lycra swim cap was placed on the head to allow for the marking of stimulation locations. We recorded motor-evoked potentials (MEPs) from the first dorsal interosseous muscle bilaterally using 4 mm Ag/AgCl electrodes placed on the muscle in a belly-tendon arrangement. A ground electrode was placed on the right wrist. MEP data were recorded and digitized at 2000 Hz using Biopac hardware and AcqKnowledge software (BIOPAC Systems Inc., Goleta, CA). Our system was carefully shielded by twisting the electrode wires, and all leads were secured in place so that they were not in contact with one another. A Magstim 70 mm figure of eight coil with a Magstim Rapid stimulator (Magstim Company Ltd, Wales, UK) was used for TMS stimulation. We localized the motor hot spot for the first dorsal interosseous muscle for each hand by stimulating at a supra-threshold level of intensity. Resting motor threshold was then determined to the nearest two percent of stimulator output that elicited MEPs of at least 50 µV on three out of six consecutive stimulations while the target muscle was at rest (cf. Triggs et al, 1994, 1999; Bernard et al., 2011). We utilized a 6 x 6 grid of points 12 mm apart for motor mapping in each hemisphere, located to encompass the majority of the motor cortical hand representation. Grid location was determined based on anatomical landmarks such that the top of the grid was placed 2 cm infero-laterally from point Cz (Sparing et al., 2008; Bernard et al., 2011). The grid extended 4.8 cm in the anterior direction and 2.4 cm in the posterior direction (Figure 1). Each site was stimulated 6 times at 110% of motor threshold with at least 6 seconds between stimulations. Stimulation was performed while the first dorsal interosseous was at rest, as determined with EMG monitoring. Mapping was
performed in both the left (dominant) and right (non-dominant) hemispheres in a counterbalanced fashion across participants.

[Figure 1 about here]

2.3 Data Processing

EMG data were filtered online and digitized (10 and 500 Hz bandwidth filtering) using a Biopac MP100 system with EMG 100C amplifiers and AcqKnowledge software (BIOPAC Systems Inc., Goleta, CA). MEP onset latency was calculated as the point at which the MEP amplitude reached two standard deviations above the mean of the amplitude of the resting state muscle activity from 500 ms before and after stimulation. The peak-to-peak amplitude was also calculated. Ipsilateral MEPs were defined as those having peak-to-peak amplitude of at least 15 µVolts (Wasserman et al., 1992; Bernard et al., 2011). The difference in onset latency between contralateral and ipsilateral MEPs (for the subset of participants who exhibited ipsilateral MEPs) was also calculated. Contralateral map areas were calculated as the number of points from which a minimum of three out of six stimulations elicited suprathreshold MEPs, similar to the cutoff of 60% employed by Pascual-Leone and colleagues (1995), whereas ipsilateral map areas were calculated as the number of points from which suprathreshold MEPs were elicited.

2.4 Statistical Analyses

Statistical analyses were performed using R software (www.r-project.org/). The cognitive and motor functional assessment scores were compared across the two age groups using independent sample t-tests. The following TMS variables were analyzed using two (age group, between subjects factor) by two (hemisphere, within subjects factor) mixed model ANOVAs: number of contralateral and ipsilateral MEPs, mean peak-to-peak amplitude of contralateral and ipsilateral MEPs, mean latency of contralateral and ipsilateral MEPs, mean latency difference between contralateral and ipsilateral MEPs, and contralateral and ipsilateral map areas. One YA was excluded from analyses of ipsilateral MEP effects and representations because their number of ipsilateral MEPs was greater than 3 standard deviations from the average.
3. Results

3.1 Age Differences in Cognitive and Motor Assessments

YA performed significantly better on the MMSE than OA ($t_{(31)}=-2.11$, $p<.05$; see section 2.1 for means), but participants in both age groups were approaching ceiling performance. Table 1 shows the mean and standard deviation of scores on the laterality and motor assessments for both age groups. Importantly, there were no age group differences in laterality of dexterity as measured by Tapping Circles ($t_{(31)}=.90$, $p>.3$; see Figure 2 for the distributions), or Edinburgh Handedness Inventory scores ($t_{(31)}=1.37$, $p>.1$). There were also no age differences in laterality on the Tapping Squares test ($t_{(31)}=.54$, $p>.5$), nor were there any differences in laterality of dexterity as measured using grip strength ($t_{(30)}=1.04$, $p>.3$).

Interestingly, there was a significant age difference in laterality of dexterity when looking at the Purdue Pegboard test ($t_{(30)}=-2.57$, $p<.05$). YA also exhibited better performance on both the bilateral ($t_{(31)}=-4.83$, $p<.001$) and the assembly ($t_{(31)}=-6.37$, $p<.001$) conditions of the Purdue Pegboard test. There were no age differences in reaction time ($t_{(31)}=1.69$, $p>.1$). [Figure 2 about here] [Table 1 about here]

3.2 Motor Threshold

A 2 x 2 ANOVA (age by hemisphere) revealed a trend for higher motor threshold in OA ($F_{(1,28)}=3.34$, $p=.08$). There was no significant main effect of hemisphere on motor threshold ($F_{(1,28)}=.03$, $p>.8$) nor was there an age by hemisphere interaction ($F_{(1,28)}=.01$, $p>.9$). Given the trend for an age difference in motor threshold, additional analyses were run using motor threshold as a covariate.

3.3 Contralateral and Ipsilateral MEP Characteristics

Table 2 contains a summary of the means and standard deviations of the TMS measures. The number of contralateral and ipsilateral MEPs was quantified and compared across age groups. OA showed a significantly larger number of MEPs contralateral to the
stimulated hemisphere when controlling for motor threshold \((F_{(1,53)}=10.07, p<.01)\). There was no significant main effect of hemisphere (dominant versus nondominant) nor was there a significant age by hemisphere interaction \((F<1, p>.3\) in both cases). There was a trend indicating that OA also showed a greater number of ipsilateral MEPs \((F_{(1,52)}=3.37, p=.07)\), though again, there was no main effect of hemisphere nor an interaction \((F<1.5, p>.3\) in both cases).

There were no age differences in peak-to-peak amplitude of ipsilateral MEPs \((F<1, p>.2)\). However, there was a significant main effect of age when looking at the amplitude of contralateral MEPs \((F_{(1,57)}=6.71, p<.05;\) Figure 3) indicating that OA had larger MEPs when controlling for motor threshold, though there was no main effect of hemisphere nor an age by hemisphere interaction \((F<1, p>.5\) in all cases).

There were no age differences in peak-to-peak amplitude of ipsilateral MEPs \((F<1, p>.2)\). However, there was a significant main effect of age when looking at the amplitude of contralateral MEPs \((F_{(1,57)}=6.71, p<.05;\) Figure 3) indicating that OA had larger MEPs when controlling for motor threshold, though there was no main effect of hemisphere nor an age by hemisphere interaction \((F<1, p>.5\) in all cases).

\[\text{[Table 2 about here]}\]

\[\text{[Figure 3 about here]}\]

We also investigated the latency of both contralateral and ipsilateral MEPs, as well as the difference between the two latencies. There was a non-significant trend for an age difference in the latency of contralateral MEPs \((F_{(1,57)}=3.75, p<.06)\), though there was no main effect of hemisphere, nor an age by hemisphere interaction \((F<1.5, p>.2\) in all cases). Onset latency of contralateral MEPs trended towards being longer in OA. Ipsilateral latencies were examined in those individuals that showed ipsilateral MEP activity. There were no age, hemisphere, nor interaction effects when looking at the latencies of ipsilateral MEPs \((F<.5, p>.6\) in all cases).

### 3.4 Motor Cortical Representations

We also evaluated whether there were age differences in the extent of the motor representations. For contralateral representations, we ran an ANOVA (age by hemisphere) on the number of points in the grid which resulted in MEPs on at least 3 stimulations (50%). Supporting our hypothesis that OA would show more diffuse motor cortical representations, a significant main effect of age \((F_{(1,57)}=5.75, p<.05;\) Figure 4) indicated that OA had larger
contralateral motor cortical representations. There was no significant main effect of hemisphere, nor was there an age by hemisphere interaction (F<1.5, p>.2 in both cases). Ipsilateral motor representations were quantified by counting the number of points in the grid that resulted in any ipsilateral MEPs given that ipsilateral MEPs were relatively scarce (9.7% of all stimulations in OA, and 2.8% of all stimulations in YA). There was a significant main effect of age on ipsilateral representations with OA representations spanning a larger number of points (F(1,49)=4.72, p<.05). Main and interaction effects with hemisphere were not significant (F<1.5, p>.2 in both instances). Figure 5 presents the contralateral and ipsilateral motor representations for representative young and older adult participants.

Finally, to investigate whether or not the extent of contralateral representations serves a compensatory mechanism or rather is reflective of dedifferentiation, we evaluated correlations with reaction time measures using Spearman’s rank correlation. In OA, there was a significant relationship indicating that a larger dominant contralateral representation was associated with higher reaction times (rho=.59, p<.05, Figure 6a) while in YA there was no significant relationship (rho=-.10, p>.6). Furthermore, the correlation coefficients were significantly different between the two age groups (Z=2.02, p<.05). A similar pattern is seen when we use a more lenient measure of contralateral representations including all points showing MEP activity (as opposed to on a minimum of 3/6 trials; OA: rho=.55, p<.05; YA: rho=-.02, p>.9; Z=1.66, p<.1; Figure 6b). Thus, it seems that the more spatially extensive contralateral representations in the dominant hemisphere of OA may be a form of less distinctive motor representations, resulting in prolonged reaction times.

4. Discussion
We found that OA have more spatially extensive contralateral and ipsilateral motor cortical representations than YA. Furthermore, the extent of contralateral representations in OA is associated with longer reaction times indicating dedifferentiation. Additionally, the characteristics of the motor evoked potentials themselves differ across the two age groups. These conclusions are supported by the following results: 1) a greater number of contralateral MEPs elicited in OA and a trend for a greater number of ipsilateral MEPs in this group; 2) larger spatial extent of contralateral and ipsilateral MEPs in OA; and 3) a larger amplitude and a trend indicating longer latency for contralateral MEPs in OA. Additionally, we observed a trend for OA to exhibit higher motor thresholds.

OA show a pattern of decreased distinctiveness in their cortical representations (Li & Lindenberger, 1999). This is particularly evident in studies focusing on the sensory systems (Li et al., 2001; Li & Sickstrom, 2002). For example, Park and colleagues (2004) have shown less specialized neural representations in the visual cortex for different categories of visual stimuli, and this finding has been recently replicated (Carp et al., In Press). Here dedifferentiation more specifically refers to when this decreased distinctiveness results in performance deficits. This is contrasted with compensation. Over-activation in prefrontal regions that is associated with better performance for cognitive tasks in OA suggests that it serves a compensatory function (Cabeza et al., 2002; Reuter-Lorenz et al., 2000; Reuter-Lorenz & Lustig, 2005).

The literature is mixed as to whether or not motor system over-activation is compensatory for OA. While some studies have shown that increased bilateral activation is associated with better task performance for older individuals, indicative of compensation (Mattay et al., 2002; for a review see Ward, 2006), there is also evidence to the contrary. For example, we recently demonstrated that activity in the primary motor cortex ipsilateral to the performing hand was correlated with higher reaction times for OA (Langan et al., 2010), supporting a dedifferentiation interpretation. The current finding of larger motor cortical representations (both contralateral and ipsilateral) in OA at rest indicates that there is a fundamental age difference in
motor cortical organization such that there may be decreased distinctiveness in the motor cortex, perhaps due to decreased intracortical inhibition reported in OA (Peinemann et al., 2001). Additionally, we found that larger contralateral representations in the dominant (left) hemisphere were associated with higher reaction times for OA. This further supports the notion of age-related dedifferentiation in the motor cortex, which may contribute to age deficits in fine motor control. Though our results indicate dedifferentiation as evidenced by a decrease in the distinctiveness of motor cortical representations and concomitant behavioral deficits, to more conclusively support this it would be necessary to map other digits within the motor cortex.

The more extensive motor cortical representations may also be due in part to age-related changes in muscle architecture (Andersen, 2003; Fling et al., 2009). With age, larger motoneurons are lost, and muscle fibers are often reinnervated by smaller neighboring motoneurons resulting in larger motor units in OA (Fling et al., 2009). Thus, this peripheral reorganization may also explain the spreading of the first dorsal interosseous representation that we observed. A similar mechanism may underlie the reduced distinctiveness of visual cortical representations in older adults. Work by Eliasieh and colleagues (2007) has demonstrated that the human retina undergoes dendritic reorganization in OA, particularly in rod, cone, and horizontal cells, though the thickness of the retina decreases with age (Polidori et al., 1993). It is therefore possible that the decrease in distinctiveness reported in the visual system is also due in part to peripheral reorganization.

A change in the architecture of motor units in OA may also underlie our findings of a larger number of MEPs, and higher contralateral MEP amplitude in OA. While it is possible that this is due to an overall increase in motor cortical excitability, it is unlikely given that we note a trend for higher motor threshold in OA. Increased excitability would predict a lower motor threshold. Though motor unit architecture changes with age (Andersen, 2003; Fling et al., 2009), OA still appear to recruit motor units in accord with the size principle (Olson et al., 1968). Thus, motor units recruited at similar thresholds are likely to be significantly larger in OA. These
larger motor units may underlie the greater MEP amplitude we see in OA (Fling et al., 2009). Furthermore, age-related reinnervation may also underlie the larger number of MEPs seen in OA. Because superficial motor fibers are more likely to be reinnervated due to the loss of larger motoneurons, more MEPs may result due to recruitment according to the size principle. Activity in these more superficial fibers is likely to be picked up with surface EMG, thus resulting in the larger number of MEPs seen in OA.

Increased recruitment of the ipsilateral motor cortex during motor task performance has been related to task difficulty (Verstynen et al., 2005; Seidler et al., 2004). Thus, increased recruitment of the ipsilateral motor cortex seen for OA (Mattay et al., 2002; Naccarato et al., 2006; Ward & Frackowiak, 2003) may be due to increased task demands. What is a simple task for YA may be more complex for OA resulting in modulation of brain activity. However, the current findings argue against this interpretation. That is, we observed more extensive motor cortical representations, particularly ipsilaterally, when participants were at rest.

Ipsilateral MEPs are not typically seen at rest in YA (Bawa et al., 2004; Carr et al., 1994; Netz et al., 1997; Ziemann et al., 1999), though we have previously shown that their presence is related to degree of handedness (Bernard et al., 2011). Thus, the mechanisms underlying the increased ipsilateral motor cortical representations, and similarly the trend indicating an overall larger number of ipsilateral MEPs in OA is of interest. Talelli et al. (2008a, 2008b) have demonstrated that with increasing age there is a decrease in interhemispheric inhibition during the performance of a background task. Furthermore, this task-related decrease in inhibition has been shown to correlate with increased ipsilateral brain recruitment during motor task performance (Talelli et al., 2008a). Perhaps then, the trend for increased ipsilateral MEPs and the corresponding more extensive ipsilateral motor representations that we observed in OA are due to differences in interhemispheric inhibition. Future studies would benefit from a direct test of the relationship between interhemispheric inhibition as measured using paired-pulse TMS and the extent of ipsilateral motor cortical representations.
Additionally, it is also important to consider the possibility that the greater amount of ipsilateral activity in OA arises due to differences in corticospinal tract organization. That is, the level of the central nervous system at which ipsilateral MEPs are generated is unclear. While callosal mechanisms as described above have been proposed (Dassonville et al., 1997), the majority of the literature on ipsilateral MEPs suggests that they are due to ipsilateral projections of the corticospinal tract (Jung & Ziemann, 1996; Ziemann et al., 1999), though these projections primarily innervate proximal muscles (Brinkman & Kuypers, 1973). Thus, while there are documented changes in interhemispheric inhibition with age, which could result in an increase in ipsilateral motor activity due to TMS stimulation, there may also be changes in the ipsilateral corticospinal tract resulting in this phenomenon. Because we did not investigate this directly, it remains an open question as to what mechanism causes ipsilateral MEPs in OA.

Characterizing the resting organization of the motor cortex in OA is important for rehabilitation interventions. TMS mapping is often used to assess damage following stroke (Liepert et al., 2000b) as well as the underlying mechanisms of treatment interventions (Liepert et al., 2000a, 2000b; Cramer & Bastings, 2000; Wittenberg et al., 2003; for a review see Hallett, 2003). However, the patients themselves often serve as their own controls, and little is known about the baseline characteristics of motor representations measured with TMS in OA. Assessing damage is therefore difficult without an appropriate point of comparison. Our analyses of the extent of motor cortical maps and number of MEPs, in addition to the characteristics of both the contralateral and ipsilateral MEPs, will provide an important reference for those assessing rehabilitation following stroke. This also may prove to be a useful point of reference for other studies looking at neurological diseases typically associated with aging, such as Parkinson’s disease.

5. Conclusions

We found that OA exhibit more diffuse motor cortical representations in the hemisphere both contralateral and ipsilateral to the recorded MEP, with larger representations correlated
with higher reaction times. This indicates that increased functional activity seen during task performance in OA may not be due to task difficulty per se, but rather may be due to differences in the resting organization of the motor cortex and/or peripheral motor unit reorganization resulting in less distinctive cortical representations. Furthermore, the greater ipsilateral representations are consistent with findings of decreased interhemispheric inhibition with age.

Acknowledgements
The authors wish to thank A. Bosma, S. Chung, S. Gluhm, N. Goodman, T. Lodgson, and R. Trivedi for help with data collection and processing. J.A.B. is supported by NIA T32 AG000114 (R. Miller, PI). Both authors are members of The LIFE Course: Evolutionary and Ontogenetic Dynamics.

Disclosure Statement.
There are no conflicts of interest. This research was approved by the University of Michigan Institutional Review Board.
References


Table 1. Mean (± SD) laterality indices (LI) and scores on motor assessments. Significant age differences are indicated with an *.

<table>
<thead>
<tr>
<th>Laterality &amp; Motor Assessments</th>
<th>YA</th>
<th>OA</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Laterality Assessments (LI)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Edinburgh Handedness Inventory</td>
<td>.72 (.16)</td>
<td>.81 (.22)</td>
</tr>
<tr>
<td>Tapping Circles</td>
<td>.13 (.06)</td>
<td>.14 (.05)</td>
</tr>
<tr>
<td>Tapping Squares</td>
<td>.09 (.05)</td>
<td>.11 (.07)</td>
</tr>
<tr>
<td>Grip Strength</td>
<td>.03 (.05)</td>
<td>.05 (.05)</td>
</tr>
<tr>
<td>Purdue Pegboard*</td>
<td>.06 (.03)</td>
<td>.02 (.05)</td>
</tr>
<tr>
<td><strong>Motor Assessments</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bilateral Pegboard (Number of Pairs)*</td>
<td>12.33 (1.62)</td>
<td>9.72 (1.48)</td>
</tr>
<tr>
<td>Pegboard Assembly (Number of Pieces)*</td>
<td>20.83 (3.49)</td>
<td>13.70 (2.92)</td>
</tr>
<tr>
<td>Dominant Hand Reaction Time (ms)</td>
<td>253.05 (56.16)</td>
<td>284.49 (50.63)</td>
</tr>
</tbody>
</table>
Table 2. Mean (±SD) of TMS measures. An * by the metric indicates a significant main effect of age.

<table>
<thead>
<tr>
<th>TMS Measures</th>
<th>YA</th>
<th>OA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>Motor Threshold (% Stimulator Output)</strong></td>
<td></td>
</tr>
<tr>
<td>Dominant Hemisphere</td>
<td>57.60 (4.91)</td>
<td>62.75 (11.83)</td>
</tr>
<tr>
<td>Non-Dominant Hemisphere</td>
<td>57.67 (5.72)</td>
<td>63 (9.00)</td>
</tr>
<tr>
<td></td>
<td><strong>Contralateral MEP Amplitude (µVolts)</strong>*</td>
<td></td>
</tr>
<tr>
<td>Dominant Hemisphere</td>
<td>136.75 (72.79)</td>
<td>209.09 (135.06)</td>
</tr>
<tr>
<td>Non-Dominant Hemisphere</td>
<td>132.72 (93.45)</td>
<td>166.48 (178.80)</td>
</tr>
<tr>
<td></td>
<td><strong>Ipsilateral MEP Amplitude (µVolts)</strong></td>
<td></td>
</tr>
<tr>
<td>Dominant Hemisphere</td>
<td>28.86 (13.54)</td>
<td>37.86 (17.57)</td>
</tr>
<tr>
<td>Non-Dominant Hemisphere</td>
<td>29.37 (18.79)</td>
<td>35.41 (26.08)</td>
</tr>
<tr>
<td></td>
<td><strong>Contralateral MEP Latency (ms)</strong></td>
<td></td>
</tr>
<tr>
<td>Dominant Hemisphere</td>
<td>21.44 (3.11)</td>
<td>20.61 (3.25)</td>
</tr>
<tr>
<td>Non-Dominant Hemisphere</td>
<td>24.13 (3.99)</td>
<td>22.64 (4.47)</td>
</tr>
<tr>
<td></td>
<td><strong>Ipsilateral MEP Latency (ms)</strong></td>
<td></td>
</tr>
<tr>
<td>Dominant Hemisphere</td>
<td>23.59 (7.02)</td>
<td>24.57 (8.44)</td>
</tr>
<tr>
<td>Non-Dominant Hemisphere</td>
<td>25.40 (12.15)</td>
<td>24.71 (7.90)</td>
</tr>
</tbody>
</table>
Figure Captions

Figure 1. Grid placement for the TMS mapping procedure. All points were 12 mm apart and placement was based upon anatomical landmarks. Mapping was done bilaterally.

Figure 2. Distribution of laterality indices for YA and OA on the Tapping Circles measure of laterality. All participants were right-handed based on self-report.

Figure 3. Average (± SD) contralateral (left) and ipsilateral (right) MEP amplitude. Contralateral MEP amplitude shows a significant main effect of age wherein OA have larger amplitude MEPs (while controlling for motor threshold; $F_{(1,57)}=6.71$, $p<.05$), *p<.05.

Figure 4. Average (± SD) number of points with MEPs on at least 50% of stimulations making up the contralateral representations (left) and average number of points with MEP activity making up ipsilateral representations (right). Significant main effects of age for both contralateral ($F_{(1,57)}=5.75$, $p<.05$) and ipsilateral ($F_{(1,49)}=4.72$, $p<.05$) representations indicate that OA have larger motor representations, *p<.05.

Figure 5. Representative contralateral and ipsilateral motor maps for a YA and an OA (MEP amplitude indicated by color, blue to red in ascending order). Both individuals are female with scores of 1.00 on the Edinburgh Handedness Inventory. The contralateral and ipsilateral maps for the left hemisphere placed on the left, and likewise for the right hemisphere.

Figure 6. A. Correlation between the contralateral representation and reaction time (RT) of the dominant hand in both YA and OA. There is a significant correlation in the OA ($\rho=.59$, $p<.05$).
but not in the YA (rho=-.10, p>.6). B. Correlations in OA (rho=.55, p<.05) and YA (rho = -.02, p>.9) when using a more lenient definition of contralateral representations.
Figure 2

The bar chart shows the distribution of participants across different tapping circles laterality index ranges. The index ranges are 0-5, 5-10, 10-15, 15-20, and 20-25.

- **YA** (black bars):
  - 0-5: 4 participants
  - 5-10: 6 participants
  - 10-15: 3 participants
  - 15-20: 3 participants
  - 20-25: 3 participants

- **OA** (gray bars):
  - 0-5: 4 participants
  - 5-10: 4 participants
  - 10-15: 5 participants
  - 15-20: 5 participants
  - 20-25: 3 participants

The chart indicates a higher number of participants in the 5-10 and 15-20 ranges for both YA and OA categories.
Figure 3

The figure shows a bar graph comparing MEP Amplitude (μVolts) between contralateral and ipsilateral hemispheres. The graph includes data for two groups: YA (black dots) and OA (gray dots). The x-axis represents the type of hemisphere (Dominant, Non-Dominant, Dominant, Non-Dominant), while the y-axis represents the MEP Amplitude (μVolts) ranging from -30 to 360 μVolts. The contralateral hemispheres for both groups show a similar MEP amplitude distribution, with a notable peak at around 210 μVolts for the YA group. The ipsilateral hemispheres show a lower amplitude distribution, with a significant asterisk (*) indicating a difference between the two groups. The error bars represent the standard deviation.
Figure 4

Motor Cortical Representation (Number of Points)

Contralateral

Ipsilateral

Hemisphere

Dominant

Non-Dominant

Dominant

Non-Dominant

 YA

OA
Figure 5

Dominant Hemisphere Stimulation

Contralateral  Ipsilateral

Non-Dominant Hemisphere Stimulation

Contralateral  Ipsilateral

 YA

 OA