The Neural Basis of Age-Related Changes in Motor Imagery of Gait: An fMRI Study

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Background. Aging is often associated with modifications of gait. Recent studies have revealed a strong relationship between gait and executive functions in healthy and pathological aging. We hypothesized that modification of gait due to aging may be related to changes in frontal lobe function.

Methods. Fourteen younger (27.0±3.6 years) and 14 older healthy adults (66.0±3.5 years) performed a motor imagery task of gait as well as a matched visual imagery task. Task difficulty was modulated to investigate differential activation for precise control of gait. Task performance was assessed by recording motor imagery latencies, eye movements, and electromyography during functional magnetic resonance imaging scanning.

Results. Our results showed that both healthy older and young adults recruited a network of brain regions comprising the bilateral supplementary motor cortex and primary motor cortex, right prefrontal cortex, and cerebellum, during motor imagery of gait. We observed an age-related increase in brain activity in the right supplementary motor area (BA6), the right orbitofrontal cortex (BA11), and the left dorsolateral frontal cortex (BA10). Activity in the left hippocampus was significantly modulated by task difficulty in the elderly participants. Executive functioning correlated with magnitude of increases in right primary motor cortex (BA4) during the motor imagery task.

Conclusions. Besides demonstrating a general overlap in brain regions recruited in young and older participants, this study shows age-related changes in cerebral activation during mental imagery of gait. Our results underscore the importance of executive function (dorsolateral frontal cortex) and spatial navigation or memory function (hippocampus) in gait control in elderly individuals.

Key Words: Mental imagery—Aging—Gait—Neuroimaging.

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Gait disorders are very common in the elderly individuals; it is estimated that only 20% of very old individuals walk normally (1–3). Specific age-related changes in spatial and temporal gait parameters have been associated with the risk of falling (4). For example, increased stride time variability predicts incidental falls in older adults. Aging can also result in increased step width variability or stride length variability, highlighting a disturbance in automatic and rhythmic motor control (5). Several neurophysiological factors may play a role in the modulation of gait in aging, from regional muscular atrophy (6) through to the degeneration of particular neurotransmitter systems, like the cholinergic system (7) or dopaminergic system (8). The presence of a gait disorder can be a marker of the future development of disease, like dementia, pathologies disturbing the frontal lobe or the connections between subcortical regions and frontal cortices (9). The frontal lobes are closely related to executive functioning, but also appear crucial for the control of gait (10). Gait disturbances may thus be indicative of subtle preclinical changes in higher order cognitive functioning.

In recent years, this relationship between executive functioning and gait control has received considerable attention (10). Impaired executive functioning has been proposed to underlie gait changes in healthy aging (11), but also in Alzheimer’s
disease (12), vascular dementia (13), frontotemporal dementia (14), and Parkinson’s disease (15). The role of executive functioning in gait control becomes apparent especially in demanding situations such as dual tasking (ie, walking while performing a cognitive task) or walking on uneven ground (10).

The use of functional neuroimaging techniques has greatly contributed to the understanding of cerebral control of gait in humans. Near-infrared spectroscopy (16), transcranial magnetic stimulation (17), nuclear neuroimaging techniques (18), and more recently electroencephalography (19) allow for direct measures of cerebral activity during actual gait performance. Functional magnetic resonance imaging (fMRI) and positron emission tomography have also been employed to study neural processes associated with gait control, through the use of motor imagery (MI) paradigms. This approach exploits the documented overlap in cerebral regions recruited during mental simulation of a movement and during actual execution of the same movement (20,21). Together, these studies have identified a network associated with MI of gait, comprising bilateral SMA, superior parietal lobules, hippocampus, parahippocampal gyri, pedunculopontine nuclei, and cerebellum (22,23).

The effects of aging on the neural substrate of gait control merits more systematic investigations given the high frequency of gait disturbances in the elderly individuals and its significance for future disease. The modification of quantitative gait parameters and the implication of cognitive functions in gait control in the elderly individuals suggest a specific pattern of brain activation during MI of gait in comparison with young adults. Our aim was therefore to compare brain function in a group of young participants and a group of healthy older adults using fMRI and a validated gait imagery task.

**METHODS**

**Participants**

Fourteen healthy young participants (4 men/10 women, mean age: 27.0 ± 3.6 years, education: 16.2 ± 2.8 years) and 14 healthy old participants (4 men/10 women, mean age: 66.0 ± 3.5 years, education: 14.6 ± 2.5 years) took part in this study. They were recruited from the community through word of mouth and public advertising. The study was approved by the local ethics committee. All participants gave written informed consent. All participants were right handed, as verified with the Edinburgh Handedness Inventory (EHI, Oldfield, 1971). Young participants obtained a mean score of 85.3 ± 17.6 and elderly participants of 87.1 ± 14.4 (p = nonsignificant). All participants had normal or corrected-to-normal vision and no neurological or orthopedic disturbances. Cognitive functioning of all elderly participants was assessed using a detailed neuropsychological test battery (Table 1) and the young participants were screened for any neurological or medical conditions interfering with brain functioning. All the older participants presented normal performances in the neuropsychological assessment.

**Table 1.** Demographic Data and Neuropsychological Test Scores of the 14 Healthy Older Participants

<table>
<thead>
<tr>
<th></th>
<th>EC</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sociodemographic</strong></td>
<td></td>
</tr>
<tr>
<td>Female/male</td>
<td>10/4</td>
</tr>
<tr>
<td>Age</td>
<td>66.0 ± 3.5</td>
</tr>
<tr>
<td>Education</td>
<td>14.6 ± 2.5</td>
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<tr>
<td><strong>Global measures</strong></td>
<td></td>
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<tr>
<td>MMSE</td>
<td>29.1 ± 0.9</td>
</tr>
<tr>
<td>Mattis</td>
<td>142.9 ± 1.2</td>
</tr>
<tr>
<td><strong>Memory</strong></td>
<td></td>
</tr>
<tr>
<td>Buschke—48 items</td>
<td>26.9 ± 3.9</td>
</tr>
<tr>
<td>Doors Test A</td>
<td>11.2 ± 1.1</td>
</tr>
<tr>
<td>Doors Test B</td>
<td>8.6 ± 1.7</td>
</tr>
<tr>
<td>Digit span forwards</td>
<td>5.9 ± 0.7</td>
</tr>
<tr>
<td>Attention/executive</td>
<td></td>
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<tr>
<td>Stroop Test</td>
<td>112.3 ± 16.3</td>
</tr>
<tr>
<td>Trail-Making Test A</td>
<td>34.6 ± 5.9</td>
</tr>
<tr>
<td>Trail-Making Test B</td>
<td>76.1 ± 20.7</td>
</tr>
<tr>
<td>Digit span backwards</td>
<td>4.2 ± 0.7</td>
</tr>
<tr>
<td><strong>Language</strong></td>
<td></td>
</tr>
<tr>
<td>Boston Naming A*</td>
<td>31.3 ± 1.9</td>
</tr>
<tr>
<td>Boston Naming C*</td>
<td>20.0 ± 0.0</td>
</tr>
<tr>
<td>Verbal fluency (category)</td>
<td>33.9 ± 7.4</td>
</tr>
<tr>
<td>Verbal fluency (letter)</td>
<td>25.0 ± 6.4</td>
</tr>
<tr>
<td>Depression/anxiety</td>
<td></td>
</tr>
<tr>
<td>HAD (depression)</td>
<td>1.4 ± 1.2</td>
</tr>
<tr>
<td>HAD (anxiety)</td>
<td>4.4 ± 2.7</td>
</tr>
</tbody>
</table>

Notes: EC = elderly controls; HAD = Hospital Depression and Anxiety scale.

*Version A of the Boston Naming test is for the age group <70 years and version C for ≥70 years.

**Motor Performance and Chronometry Ability**

To assess individual participant’s motor performance and chronometry ability (CA), we administered two mental chronometry tests, both evaluating performed and imagined times of an action. The first was a version of the Timed “Up and Go” (TUG) test developed by Beauchet and coworkers (24). The initiation, acceleration, deceleration, preparation of turning, and turning component of the TUG are also involved in the “difficult” condition of our fMRI task. The second was the Ten Meter test. Participants completed two trials on both the real and imagined version of both tasks. On the imagined TUG, the experimenter gave the cue to start imagining, and the participant was instructed to pronounce the word “stop” when they had completed the mental task. The Ten Meter test was similar in design to the TUG, but assessed CA while walking in a straight line, similar to the fMRI task (which also involved walking straight for 10 m). All participants performed two trials of the Ten Meter and the TUG and the mean of these two trials were used as outcome.

Individual CA scores were calculated on the basis of the delta time scores on the TUG and Ten Meter test. Delta times were calculated according to the following formula: 

\[ \delta t = \frac{(T_{real} - T_{imag})(T_{real} + T_{imag})}{2} \]

CA score was calculated as the difference between the actual test time and the imagined time. The delta time scores were calculated as the difference between the real and imagined times on the TUG and Ten Meter test.

Delta time scores on the TUG and Ten Meter test. Delta time scores were calculated as the difference between the real and imagined times on the TUG and Ten Meter test.
taken as the positive mean of the two delta times for each participant. The lower the CA score, the smaller the difference between real and imagined performance times, and therefore the higher CA.

fMRI Procedure

Materials.—Participants were scanned during a single fMRI session lasting about 30 minutes. All functional and structural MRI data were acquired on a 3T whole body MRI system (Trio TIM, Siemens, Germany) with a 12-channel head coil at the Brain & Behaviour Laboratory (BBL) of the University of Geneva. Visual stimuli were presented on a projection screen inside the scanner using E-prime (E-prime 1.0, Psychology Software Tools Inc., Pittsburgh, PA). Gaze direction and pupil diameter were logged with an MRI compatible long range eye tracker (EyeTrac 6, Applied Science Laboratories).

Scanning protocol.—Whole-brain functional images were collected using a susceptibility-weighted EPI sequence (TR/TE = 2,100/30 ms; flip angle = 80°; PAT factor = 2; FOV = 205 mm; matrix size = 64 × 64 pixels). Thirty-six transversal slices were acquired sequentially with a 3.2 mm thickness and an interslice gap of 0.64 mm, yielding a voxel size of 3.2 mm isotropic. High-resolution whole-brain anatomical scans were acquired with a T1-weighted, 3D sequence (MPRAGE; TR/TI/TE = 1,900/900/2.27 ms; flip angle = 9°; voxel dimensions = 1 mm isotropic; 256 × 256 × 192 voxels).

fMRI activation task.—Stimuli for the fMRI task consisted of 12 photos, 6 each of 2 different outdoor walkways, located in the garden of the rehabilitation unit of the Geneva University Hospital, each taken from a slightly different perspective, side or angle (Figure 1) that have been described previously (25). Both walkways were about 1 m wide. One walkway had a smooth tarred surface (providing stimuli for the “easy” condition), whereas the other had a surface consisting of cobble stones of between 4 and 30 cm in diameter (“difficult” condition). Participants performed three different tasks on the two kinds of walkways: MI, visual imagery (VI), and a perceptual control (C) condition.

In the MI task, participants were shown the photos of the different walkways with two horizontal lines drawn across them: a grey line at the bottom to indicate the start of the trajectory and a black line at a distance of 10 m (as measured on the actual path). Participants were asked to start imagine walking from the grey line to the black line as soon as the picture appeared. They had to press a button when they imagined that they had reached the red line on the walkway. Focusing on the kinesthetic component of MI, they were instructed to imagine walking as vividly as possible, as if feeling their limbs moving (but without actually moving them), and from a first-person perspective.

In the VI task, participants were presented with the same photos as in the MI task, but this time a black disk (25 cm in diameter) was shown on each photo at the start of the trajectory (on the grey line). Participants were asked to imagine this disk moving from the blue to the red line at a constant speed, similar to walking speed. They pressed a button when they had imagined the disk to arrive at the red line.

In the control task, participants were presented with the same pictures as in the MI task, but in this case, the lines on the walkways were either both grey or both black. Participants were instructed to simply inspect each photograph for a fixed period of time (6 seconds in half of the trials and 10 seconds in the other half). After offset of the picture, participants were cued to press either one button if the lines on the walkway were black or another in case they were blue.

Procedure.—The three different tasks were presented in a block-wise fashion, with two blocks per task, and therefore six blocks in total, that have been described previously (25). All participants received detailed instructions and underwent a training session on all three conditions of the task before the scanning session, until they were confident that they were able to perform the task at the best of their capacities.

Behavioral Data Analysis

For each participant, the mean imagery time for both MI and VI was calculated by averaging the trials for each condition. Imagery time was taken as the time between the onset of the picture and the button press. We used an eye-tracker monitor to check online that all participants kept their eyes open and adhered to the actual task instructions (ie, moved their eyes along the walkways). Moreover, following the procedure by Bakker and coworkers (22), we also verified task adherence by investigating the effect of path surface (ie, task difficulty) for the MI and the VI tasks. As the black disk is not impeded by path surface, task difficulty should have a larger effect on MI than on VI. Therefore, a 2 × 2 repeated-measures analysis of variance (ANOVA) of imagery times was carried out, with task (MI vs VI) and difficulty (easy vs difficult) as within-participant variables.

EMG Recording/Analysis

During the fMRI session, muscle activity of the right leg was recorded to control for overt muscle movements. Muscle activity (EMG) was recorded with a modular data acquisition system (MP150, BIOPAC Systems Inc.), with a sampling rate of 10,000 Hz. A pair of carbon-wired MRI compatible electrodes were placed 5 cm apart along the muscle bellies of the right tibialis anterior. The reference electrode was placed on the center of the patella. Details of EMG analysis have been previously described (25).
Functional Data Analysis

First-level analysis.—Functional images were preprocessed and analyzed using standard methods implemented in SPM5 (Wellcome Department of Imaging Neuroscience, London), which have been described previously (25). The experimental conditions were Motor Imagery—Easy (MIE) Motor Imagery—Difficult (MID) Visual Imagery—Easy (VIE) Visual Imagery—Difficult (VID) and Control (C) (Figure 1).

Second-level analysis.—Contrast images obtained in the individual analyses were entered into one flexible factorial design, which included group and condition as main factors, as well as their interaction. The main contrasts of interest were MI > C, probing for activation associated with MI, and MID > MIE, investigating the effect of path surface (ie, task difficulty). MI > VI was also included to investigate the motor specificity of the effects. Imagination times were added as a linear parametric nuisance covariate to make sure that any difference between conditions or groups was not due simply to a difference in time taken to perform the task. CA scores were also added as a second covariate to control for differences between groups in CA. Statistical analyses were performed on a voxel-wise basis across the whole brain, all thresholded at $p < .001$ (uncorrected) with a minimum cluster size of 20 voxels, unless otherwise specified.
Anatomical labels of activated clusters were determined using the xjView toolbox (http://www.alivelearn.net/xjview) and the Anatomy Toolbox version 1.5.

Regression.—To explore whether there were any areas in which activity during gait imagery correlated with executive functioning and according to the previous observations in matter of association between cognitive function and gait (25), we added the Stroop interference score as a linear regressor to the MI > C contrast of our task using the multiple regression option in SPM5. The Stroop interference score refers to the difference in response times related to the suppression of word reading in favor of color naming, which assesses inhibitory control. Walking requires the quick online processing of a flow of information, to adapt motor actions to the environmental conditions and inhibiting the nonrelevant information, which is crucial for gait.

Results

Behavioral Results

Motor performance and CA.—Mean performed (\(T_{\text{real}}\)) and imagined times (\(T_{\text{imag}}\)) of the two mental chronometry tests, as well as CA scores, are presented in Table 2. From the table, it can be seen that elderly participants took longer to perform both tasks (significant only for the TUG), indicating a slight reduction in walking speed.

Both groups always imagined doing the tasks faster than they executed them in reality. CA scores differed significantly between groups (\(F(1,13) = 7.21, p < .012\)), indicating that CA was slightly worse in the elderly participants.

fMRI gait imagery task.—Mean imagery latencies during the different conditions of our fMRI task are summarized in Table 3. Responses in the control condition (deciding on the color of the lines) were 100% correct for all participants. A repeated-measure ANOVA with task (MI and VI) and condition (easy and difficult) as within-participants factors revealed that for the young participants, there was a main effect of condition \((F(1,13) = 40.64, p < .001)\), and an interaction between task and condition \((F(1,13) = 9.40, p < .01)\), but no main effect of task \((F(1,13) = 1.11, p = \text{nonsignificant})\). This indicates that they took longer to perform the difficult than the easy condition for both MI and VI and that the difference between easy and difficult was significantly greater for the MI than for the VI task. For the old participants, there was only a main effect of condition \((F(1,13) = 79.63, p < .001)\), but no main effect of task \((F(1,13) = 0.23, p = \text{nonsignificant})\) and no interaction \((F(1,13) = 0.38, p = \text{nonsignificant})\). Therefore, elderly participants showed an equal difference in imagery times between the easy and difficult condition for both the MI and VI tasks. When the two participant groups were collapsed, we found a main effect of condition \((F(1,27) = 106.95, p < .001)\) and an interaction between tasks and condition \((F(1,27) = 5.59, p < .05)\).

EMG Results

The mean normalized muscle activity values for MI and VI are presented in Table 4. A repeated-measures ANOVA with task as within-participants factor revealed no difference in EMG values between MI and VI for the young participants \((F(1,13) = 1.47, p = \text{nonsignificant})\) nor for the elderly participants \((F(1,13) = 0.87, p = \text{nonsignificant})\). There were also no differences between groups for MI \((F(1,27) = 0.02, p = \text{nonsignificant})\) nor for VI \((F(1,27) = 1.34, p = \text{nonsignificant})\). This indicates that young and old participants presented the same muscle activity during MI and VI and that this muscle activity was similar in both groups.

Imaging Results

Motor imagery effect (MI > C).—For the main contrast MI > C, we found widespread activation in both groups in a network of areas including bilateral supplementary motor area (BA6) extending to midcingulate cortex, superior medial prefrontal cortex, lateral inferior BA4, right inferior prefrontal cortex (incl. BA44/45/47) extending to the insula and right postcentral gyrus, and the cerebellum. When we compared groups directly for this contrast \((\text{MI > C}_\text{elderly}) > \text{(MI > C}_\text{young})\), we found stronger activation in elderly participants in the left middle frontal gyrus (BA10), right SMA, and right superior orbitofrontal cortex (BA11). The parameter estimates from these three clusters showed a group difference only for the MI task and not for the VI task, indicating that only imagery for gait (and not imagery for visual movement) in our task is modulated by age (Figure 2 and Supplementary Table 1). There were no clusters that were activated stronger in the young than in the

| Table 2. Mean ± SD \(T_{\text{real}}, T_{\text{imag}}\) on the TUG and TM, and Chronometry Ability Scores |
|---------------------------------|-------------------------------------------------|---------------------------------|--------------------------|
| TUG | \(T_{\text{real}}\) | \(T_{\text{imag}}\) | \(T_{\text{real}}\) | \(T_{\text{imag}}\) | Chronometry Ability |
|---------------------------------|-------------------------------------------------|---------------------------------|--------------------------|
| Young participants | 8.08 ± 0.93 * | 6.67 ± 1.59 | 7.59 ± 0.83 | 6.83 ± 1.18 | 0.19 ± 0.13 * |
| Old participants | 9.33 ± 1.38 * | 6.64 ± 2.29 | 7.94 ± 0.84 | 7.50 ± 2.02 | 0.33 ± 0.14 * |

*Notes: \(T_{\text{imag}}\) = imagined performance time; \(T_{\text{real}}\) = real performance time; TM = Ten Meter; TUG = Timed Up and Go test.
*p < .05 between young and old participants.
For the elderly participants. When we compared groups for the contrast MI > VI, using the same threshold as for the contrast MI > C, we did not find any stronger activation in elderly participants.

**Correlation of activation with Stroop task.**—For the elderly participants, we investigated whether there were any regions that showed a correlation between activation during imagery of gait and behavioral performance on the Stroop test (interference condition). We found a single cluster in the right primary motor cortex (BA4), where activity was tightly and positively correlated with performance on the Stroop test (Figure 3 and Supplementary Table 2), indicating that the better executive functioning, the stronger the recruitment of the right primary motor area during MI.

**Task difficulty (MID > MIE).**—To investigate differential activation for precise control of gait, we looked at the effect of path surface during imagine walking (ie, task difficulty). In the elderly group, we found differential activation for precise control of gait, we looked at the kinesthetic modality of MI. We found greater activation in some subcortical structures, in particular the substantia nigra and putamen. In the only previous study assessing the effect of aging on the human supraspinal locomotor and postural control by fMRI, although the authors did not assess the cognitive function of their elderly participants, a prominent activation was also observed in the SMA. The correlation between activation during imagery of gait and performance on the Stroop test in the right primary motor cortex that we found reinforces the notion of a strong link between executive function and gait.

The frontal lobes are highly susceptible to age-associated changes, including disruption of frontotriatal circuits by diffuse white matter changes, gray matter atrophy, and reduction of dopamine activity in the frontal cortex. Interestingly, with regard to the greater activation in the orbitofrontal cortex in the elderly participants, previous studies indicate that elderly patients with freezing of gait present a specific hypometabolism in these orbitofrontal regions. These metabolic and morphological changes may contribute to deficits in executive functioning as commonly seen in the elderly participants.

Another main result of this study was a greater activation in the left hippocampus in elderly participants relative to young participants, specifically for the MID > MIE contrast that probed activations associated with the higher order, more precise control of gait. The hippocampus and adjacent regions are well known to be involved in age-related changes in memory and spatial navigation but have also been implicated in gait. In a positron emission tomography study on gait, Malouin and coworkers have shown the involvement of a significant recruitment of the left parahippocampal region during walking. The specific role of the hippocampus/medial temporal lobe in gait may therefore be related to topographical memory and mental navigation. Animal models also suggest that the hippocampal formation provides voluntary motor systems with continually updated feedback on environmental conditions. Previous studies on anesthetized rats revealed a strong interaction between the hippocampus and the pedoculopontine studies on gait imagery. Elderly participants recruited a network that was similar to that recruited by young participants, but with some critical differences, discussed later.

As predicted, the elderly participants exhibited significantly greater activation than the young participants during MI in two prefrontal regions (left BA10 and right BA11) and the right SMA. The given instruction focusing on the kinesthetic modality of MI could contribute to these results. Indeed, Guillot and coworkers showed that the kinesthetic modality of MI was associated with more activity in motor-associated brain regions. Remarkably, the elderly participants also showed a correlation between activation during imagery of gait and performance on the Stroop test in the right primary motor cortex (BA4). Additionally, they showed stronger activation in some subcortical structures, in particular the substantia nigra and putamen. In the only previous study assessing the effect of aging on the human supraspinal locomotor and postural control by fMRI, although the authors did not assess the cognitive function of their elderly participants, a prominent activation was also observed in the SMA. The correlation between activation during imagery of gait and performance on the Stroop test in the right primary motor cortex that we found reinforces the notion of a strong link between executive function and gait.

**Discussion**

In this study, we explored the effects of aging on the neural substrate of gait control, using a MI paradigm. We have identified a network of activation in young participants during MI of gait (MI > C), comprising bilateral primary motor cortex and SMA, right prefrontal cortex and cerebellum, which is highly consistent with previous neuroimaging studies on gait imagery. Elderly participants recruited a network that was similar to that recruited by young participants, but with some critical differences, discussed later.

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The frontal lobes are highly susceptible to age-associated changes, including disruption of frontotriatal circuits by diffuse white matter changes, gray matter atrophy (28), and reduction of dopamine activity in the frontal cortex (29). Interestingly, with regard to the greater activation in the orbitofrontal cortex in the elderly participants, previous studies indicate that elderly patients with freezing of gait present a specific hypometabolism in these orbitofrontal regions (30). These metabolic and morphological changes may contribute to deficits in executive functioning as commonly seen in the elderly participants (28,29).

Another main result of this study was a greater activation in the left hippocampus in elderly participants relative to young participants, specifically for the MID > MIE contrast that probed activations associated with the higher order, more precise control of gait. The hippocampus and adjacent regions are well known to be involved in age-related changes in memory and spatial navigation but have also been implicated in gait. In a positron emission tomography study on gait, Malouin and coworkers (23) have shown the involvement of a significant recruitment of the left parahippocampal region during walking. The specific role of the hippocampus/medial temporal lobe in gait may therefore be related to topographical memory (31) and mental navigation (32). Animal models also suggest that the hippocampal formation provides voluntary motor systems with continually updated feedback on environmental conditions (33). Previous studies on anesthetized rats revealed a strong interaction between the hippocampus and the pedoculopontine

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**Table 3. Mean ± SD Imagery Times in Seconds**

<table>
<thead>
<tr>
<th>Task</th>
<th>MI</th>
<th>VI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Condition</td>
<td>Easy</td>
<td>Difficult</td>
</tr>
<tr>
<td>Young participants</td>
<td>8.7±2.4</td>
<td>10.5±3.2</td>
</tr>
<tr>
<td>Old participants</td>
<td>9.0±3.0</td>
<td>10.9±3.8</td>
</tr>
</tbody>
</table>

Notes: MI = motor imagery; VI = visual imagery.

**Table 4. Mean ± SD EMG Values for the Motor Imagery and Visual Imagery Normalized for the Control Condition**

<table>
<thead>
<tr>
<th>Task</th>
<th>MI</th>
<th>VI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Young participants</td>
<td>1.05±0.30</td>
<td>0.99±0.28</td>
</tr>
<tr>
<td>Old participants</td>
<td>1.05±0.22</td>
<td>1.10±0.16</td>
</tr>
</tbody>
</table>

Notes: MI = motor imagery; VI = visual imagery.
Figure 2. Activation in the MI > C contrast elderly participants versus young participants.
nucleus mediated by cholinergic input (34). One speculative hypothesis is that the cholinergic pathway reduction due to aging (7) may have contributed to this increased activation in the hippocampus in the elderly participants.

These effects of aging on brain activation during cognitive tasks could be explained by the compensation hypothesis, which holds that aging is associated with a decline in brain function and performance, inducing hyperactivation of specific brain regions as a compensatory mechanism (35). The dedifferentiation hypothesis, according to which older adults tend to present a nonselective recruitment of brain regions relative to young adults (36), could be another explanation. The greater activation in the right SMA (BA6), the right orbitofrontal cortex (BA11), the left dorsolateral frontal cortex (BA10), and the left hippocampus observed in old adults reflects a difficulty in recruiting specialized neural mechanisms (37). Perhaps associated with this inability of selective task-associated cerebral recruitment, the elderly participants may have used different strategies to perform the task. Our behavioral data indicate that the young participants showed a greater modulation of imagination time by task difficulty during the MI task than during the VI task. In contrast, the elderly participants showed an equal modulation by type of path for both tasks. Performance on our task could be regarded as worse in elderly participants than in young participants, given that they fail to show an interaction between task and difficulty. Moreover, like previously observed (38), the motor CA of our elderly participants was worse than that of the young participants, as evidenced by a larger discrepancy between performed and imagined times.
on the two mental chronometry tests. Finally, the increased left hippocampal activation seen for the MID > MIE contrast in the elderly participants relative to young participants suggests that the navigational aspect of walking on cobble stones may have been more challenging to them than to young participants. Therefore, the additional cortical and subcortical areas recruited in older adults may constitute a compensatory mechanism in an attempt to achieve normal performance levels on the MI task, in the light of reduced imagery performance and ability.

As a study limitation, it should be noted that most of the activations were obtained with a standard strict threshold of \( p < .001 \) at the voxel level, with a cluster threshold of \( k > 20 \). This combination of threshold and cluster criteria selects reliable effects when changes in BOLD (Blood-oxygen-level dependent) have relatively weak amplitudes and imprecise onsets as for mental imagery or other purely internal (non-sensory) mental events (39). Importantly, all activations reported were selective and concerned areas well known to be involved in MI of gait. Furthermore, we found a significant correlation in the right primary motor cortex, prefrontal cortex, and an independent cognitive measure (Stroop test). So this suggests that our effects were reliable and task related.

**CONCLUSION**

We have demonstrated that aging results in an increased activation during imagery of gait in the prefrontal cortex (specifically, in the left MFG [BA10] and right superior OFC [BA11]) and the right SMA, with additional increases in left hippocampal activation during more precise gait control. In addition, executive functioning of our elderly participants correlated with activity during MI of gait in the right primary motor cortex. These findings emphasize the age-related modification of brain functioning and the important role of executive function in the control of gait in healthy human participants. Future studies should investigate how these neural circuits are modified in pathologies affecting gait control, such as primary progressive gait apraxia, and further help to guide and assess neurorehabilitation techniques.

**SUPPLEMENTARY MATERIAL**

Supplementary material can be found at: http://biomedgerontology.oxfordjournals.org/

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**CONFLICT OF INTEREST**

The authors report no conflicts of interest.

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