Entorhinal Cortex Volume Is Associated With Dual-Task Gait Cost Among Older Adults With MCI: Results From the Gait and Brain Study

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Abstract

Background: Low dual-task gait performance (the slowing of gait speed while performing a demanding cognitive task) is associated with low cognitive performance and an increased risk of progression to dementia in older adults with mild cognitive impairment. However, the reason for this remains unclear. This study aimed to examine the relationship between dual-task cost and regional brain volume, focusing on the hippocampus, parahippocampal gyrus, entorhinal cortex, and motor and lateral frontal cortices in older adults with mild cognitive impairment.

Methods: Forty older adults with mild cognitive impairment from the “Gait and Brain Study” were included in this study. Gait velocity was measured during single-task (ie, walking alone) and dual-task (ie, counting backwards, subtracting serial sevens, and naming animals, in addition to walking) conditions, using an electronic walkway. Regional brain volumes were derived by automated segmentation, using 3T magnetic resonance imaging.

Results: Partial rank correlation analyses demonstrated that a smaller volume of the left entorhinal cortex was associated with higher dual-task costs in counting backwards and subtracting serial sevens conditions. Subsequent logistic regression analyses demonstrated that a smaller volume of the left entorhinal cortex was independently associated with higher dual-task cost (slowing down >20% when performing cognitive task) in these two conditions. There were no other significant associations.

Conclusions: Our results show that lower dual-task gait performance is associated with volume reduction in the entorhinal cortex. Cognitive and motor dysfunction in older adults with mild cognitive impairment may reflect a shared pathogenic mechanism, and dual-task-related gait changes might be a surrogate motor marker for Alzheimer’s disease pathology.

Keywords: Dual task, Entorhinal cortex, Hippocampus, Gait, Mild cognitive impairment

Mild cognitive impairment (MCI) is typically described as a clinical state between normal cognition in aging and very early dementia. Although older adults with MCI have an increased risk of progressing to Alzheimer’s disease (AD) and other dementias, almost one third of the older adults with MCI remain clinically stable or even revert to normal (1). Early detection of the risk factors associated with progression to dementia among older adults with MCI is, therefore, necessary for the appropriate targeting of prompt treatments.

Numerous studies have shown that slow gait is a risk factor associated with the development of MCI, dementia, and a more rapid rate of cognitive decline (2,3). This may be because gait control, which relies on executive function (4,5), shares common brain networks with the cognitive processes essential for the planning and
monitoring of goal-directed behavior. Previous studies have elucidated this cognition-motor interaction using the dual-task gait paradigm (walking while performing a cognitive task) (4,6–8). These studies have revealed an association between slowing gait while performing dual tasking and lower level of cognitive function (9,10). It has been suggested that the dual-task gait paradigm examines the cognitive component of locomotion and can provide a window to the mechanism of gait control and cognitive performance (5,11). However, the longitudinal association between slowing gait while performing dual tasking and the risk of progression to dementia among older adults with MCI was, until recently, unknown.

A recent prospective study answered this question and demonstrated that a high dual-task gait cost (slowing down >20% when dual tasking compared with single tasking), which adjusts for an individual’s baseline gait velocity, was associated with a twofold to threefold increased risk of progression to dementia, whereas slow single-task gait velocity was not associated with progression to dementia (12). These results suggest that, among older adults with MCI, using solely a slow gait velocity threshold may be insufficient to identify individuals at higher risk of progression to dementia. In contrast, the dual-task paradigm could be applied as a clinical motor marker for early detection of older adults with MCI at higher risk of progression (12).

Considering the close association between high dual-task gait cost and progression to dementia, the pathways involved in these two processes may share the same neural pathology, including atrophy in the hippocampus, parahippocampal gyrus, and/or entorhinal cortex (13–17). Although the neural pathways underlying dual-task gait performance are not well established, for single-task gait, a slow gait speed has been related to specific cognition-related cerebral metabolic and structural abnormalities (18–22), and AD neuropathology (23). For example, studies using positron emission tomography showed that lower gait velocity was associated with lower glucose metabolism in the posterior cingulate cortex of older adults without a neurologic disease, which is an early sign of AD (20,21). More recently, a longitudinal study found that the association between gait slowing and cognitive impairment is supported by a shared neural substrate that includes a decreased volume of the right hippocampus (22).

Understanding the association between dual-task gait performance and regional brain volumes, vulnerable to AD pathology in older adults with MCI, may provide a neurological explanation for a high dual-task gait cost being associated with an increased risk of progression to dementia. The purpose of this study was to examine the neural substrate of dual-task gait cost, focusing on the medial temporal areas, particularly the hippocampus, parahippocampal gyrus, and entorhinal cortex, which are vulnerable regions in AD. We also introduced control regions of interest thought to be involved in motor control and dual tasking, including the motor and lateral frontal cortices, which completely overlap with the dorsolateral prefrontal cortex, and other neighboring regions such as the ventrolateral prefrontal and premotor cortices (24–26).

Methods
Participants
The Gait and Brain Study is a prospective cohort study designed to elucidate whether quantitative gait impairments can predict incident cognitive and mobility decline as well as progression to dementia among community-dwelling older adults without dementia at baseline (10). The experimental design and rationale have been described elsewhere (10,12). Participants were included in this study if they fulfilled MCI criteria and had 3T magnetic resonance imaging (MRI) of the brain performed during baseline assessment, introduced in the second wave of the “Gait and Brain Study” in 2010.

We defined MCI by a score of 0.5 on the clinical dementia rating scale and fulfillment of the following criteria: (a) subjective cognitive complaints; (b) objective cognitive impairment in at least one of the following cognitive domains: memory, executive function, attention, and language (27,28); (c) preserved daily activities, confirmed by clinician interviews; and (d) the absence of dementia, on the basis of criteria from the Diagnostic and Statistical Manual of Mental Disorders, fourth edition revised.

The exclusion criteria included diagnosis of a terminal illness, a life expectancy of less than 12 months, pending nursing home placement, hip or knee joint arthroplasty within the preceding 6 months, inability to walk 10 m independently without any gait aid, and the diagnosis of dementia.

Written informed consent was obtained from all participants prior to the study. The study was conducted in accordance with the ethical standards outlined in the Declaration of Helsinki, and the research protocol was approved by the University of Western Ontario Health Sciences Research Ethics Board (clinicaltrials.gov identifier: NCT03020381).

Assessments
Comprehensive assessments, including gait performance, medical, and cognitive evaluations, were conducted at the Gait and Brain Lab, at Parkwood Institute, and brain MRI were performed at the Centre for Functional and Metabolic Mapping at the Robarts Research Institute using a 3T Tim Trio and 3T Magnetom Prisma MRI scanners (Siemens). Both centers are affiliated to the University of Western Ontario.

Quantitative gait assessment
Gait velocity (centimeter per second) was assessed during single and dual tasks using an electronic walkway (GaitRite Systems, 600 cm long and 64 cm wide; CIR Systems, Inc., Sparta, NJ). Start and end points were marked on the floor, 1 m from both the walkway start and end point, to avoid recording the participants’ acceleration and deceleration phases.

The single-task trial consisted of walking the length of the walkway at the participant’s usual pace. For the dual-task trials, participants walked at their usual pace with no instruction to prioritize the gait or cognitive task while performing the following cognitive tasks aloud: (a) counting backwards from one hundred by ones, (b) subtracting serial sevens from one hundred, or (c) naming animals. To balance and minimize the effects of learning and fatigue, the order of the single and dual tasks was randomized. The rationale and reliability behind the use of this protocol in people with MCI have been described elsewhere (29). To examine the effect of the addition of a cognitive task on gait performance, gait velocity in dual-task condition was compared with single-task condition (ie, the dual-task gait cost) using the following formula: ([single-task gait velocity − dual-task gait velocity]/single-task gait velocity) x 100. Participants who had a dual-task cost of more than 20% were considered to have a high dual-task cost (12). The gait assessments were conducted in a quiet, well-lit room, and participants wore comfortable footwear.

MRI scanning protocol and image processing
The MRI protocol included 3D T₁-weighted magnetization-prepared rapid acquisition gradient echo images, with matrix size,
256 × 256 × 160; field of view 256 × 256 × 192 mm, repetition time = 2.3 ms, and echo time = 2.9 ms. All 3D T₁-weighted MRI scans were visually inspected for abnormalities.

Subcortical structures including the hippocampus, parahippocampal gyrus, and entorhinal cortex (Figure 1), and regions of the motor and lateral frontal cortices were demarcated using the FreeSurfer software (version 5.3.0, available at http://surfer.nmr.mgh.harvard.edu/). A composite volume of the lateral frontal cortex was calculated by adding the volumes of superior, middle, and inferior frontal gyri, excluding the frontopolar and orbitofrontal regions of each hemisphere (30).

The initial processing of the 3D T₁-weighted images included the following steps: removal of nonbrain tissue, automated Talairach transformation, segmentation of the subcortical white matter and deep gray matter volumetric structures, intensity normalization, tessellation of the gray matter/white matter boundary, automated topology correction, and surface deformation to optimally place the gray matter/white matter and gray matter/cerebrospinal fluid boundaries. Subsequently, all participants’ data were visually evaluated to ensure that the target areas were correctly segmented. If errors were detected, the participants’ data were processed to correct misidentified regions using a statistical parametric mapping and checked visually for a second time.

Medical and cognitive assessments
Participants were interviewed on relevant sociodemographic and clinical variables. Global cognitive function was assessed by the Mini–Mental State Examination (31) and the Montreal Cognitive Assessment (MoCA) test (32), which examines six domains of overall cognitive function. Both tests have a maximum score of 30 points, with higher scores indicating higher overall cognitive function. Executive function and episodic memory were evaluated using the Trail-Making Test (TMT) (33) and the Rey Auditory Verbal Learning Test (RAVLT) (34). The TMT consists of parts A and B; TMT-A assesses visual search, motor speed skills, and attention, and TMT-B evaluates working memory and task shifting.

Data Analysis
The participants’ characteristics were summarized using means and SD or frequencies and percentages, as appropriate (Table 1). To assess associations between cerebral regions (hippocampus, parahippocampal gyrus, entorhinal cortex, and motor and lateral frontal cortices of each hemisphere) and gait variables (single- and dual-task gait velocity and dual-task cost), partial correlation analyses adjusted for intracranial volume were performed. In the case of dual-task costs, partial rank correlation analyses were used for all three conditions, due to their nonparametric distribution. A Bonferroni correction was applied to avoid type 1 error because correlation analyses were repeated 10 times for each gait variable (p = .05/10 = .005). Multiple regression for parametric variables (ie, gait velocity) and logistic regression analyses for nonparametric variables (ie, dual-task costs) were then performed to examine respective associations among regional brain volumes and gait variables on the basis of all resultant significant correlation coefficients. For the logistic regression analysis, dual-task cost > 20% was used as the dependent variable (12). The regression analyses were adjusted for age, sex, intracranial volume, education level, and MoCA score. Statistical analyses were conducted using SPSS 21.0 (IBM Corporation, Chicago, IL).

Results
Although none of the participants met the exclusion criteria, two participants were excluded due to anatomical abnormalities causing mis-segmentation of target areas during image processing. Thus, a total of 40 Caucasian older adults with MCI were included in these analyses. Demographic and clinical characteristics are presented in Table 1. The mean age was 74.2 ± 6.0 years, and 42.5% of the participants were women. Performance during cognitive testing revealed that half of the participants exhibited amnestic-MCI, single and multiple domains (37.5% and 10%, respectively). Mean score of the RAVLT in the delayed recall was 6.8 out of 15. Aligned with the

Table 1. Participants’ Characteristics

<table>
<thead>
<tr>
<th>Variables</th>
<th>Participants in this Study (n = 40)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y), mean (SD)</td>
<td>74.2 (6.0)</td>
</tr>
<tr>
<td>Females, n (%)</td>
<td>17 (42.5)</td>
</tr>
<tr>
<td>Education (y), mean (SD)</td>
<td>13.3 (2.6)</td>
</tr>
<tr>
<td>Total number of medications taken</td>
<td>8.0 (4.3)</td>
</tr>
<tr>
<td>Total number of comorbidities</td>
<td>5.8 (2.8)</td>
</tr>
<tr>
<td>Previous fall, n (%)</td>
<td>8 (20.0)</td>
</tr>
<tr>
<td>GDS, mean (SD)</td>
<td>2.3 (2.2)</td>
</tr>
<tr>
<td>MMSE, mean (SD)</td>
<td>27.7 (2.4)</td>
</tr>
<tr>
<td>MoCA, mean (SD)</td>
<td>23.7 (3.7)</td>
</tr>
<tr>
<td>Rey auditory verbal learning (/15), mean (SD)</td>
<td>6.8 (2.0)</td>
</tr>
<tr>
<td>Rey auditory verbal learning (/45), mean (SD)</td>
<td>16.8 (5.1)</td>
</tr>
<tr>
<td>TMT-A (s), mean (SD)</td>
<td>44.5 (15.5)</td>
</tr>
<tr>
<td>TMT-B (s), mean (SD)</td>
<td>142.9 (151.9)</td>
</tr>
<tr>
<td>Amnestic-MCI single domain, n (%)</td>
<td>15 (37.5)</td>
</tr>
<tr>
<td>Amnestic-MCI multiple domains, n (%)</td>
<td>4 (10.0)</td>
</tr>
<tr>
<td>Gait velocity (cm/s), mean (SD)</td>
<td>Single gait 110.1 (20.5)</td>
</tr>
<tr>
<td>Counting gait</td>
<td>106.3 (26.2)</td>
</tr>
<tr>
<td>Serial sevens gait</td>
<td>94.7 (31.3)</td>
</tr>
<tr>
<td>Naming animals gait</td>
<td>96.1 (28.1)</td>
</tr>
</tbody>
</table>

Note: GDS = Geriatric Depression Scale; MCI = mild cognitive impairment; MMSE = Mini–Mental State Examination; MoCA = Montreal Cognitive Assessment; TMT = Trail-Making Test.

*Final score is the number of words remembered out of a list of 15 in trial 3 (delayed recall). †Final score is the total number of words remembered for trials 1–3.
good functionality of this cohort, the mean single-task gait velocity was 110.1 cm/s, above the accepted cutoff for impairment (35).

The results of the partial correlation analyses for regional brain volumes and gait variables are presented in Table 2. A lower volume of the left entorhinal cortex was significantly associated with slower dual-task gait velocity under all three conditions and a higher dual-task gait cost in counting backwards and subtracting serial sevens conditions (Figure 2). There were no significant associations between gait variables and other regional brain volumes.

When regression analyses were performed on all significant correlation coefficients, results remained the same: The lower left entorhinal volume was associated with slower gait velocities on all dual tasks and a higher dual-task gait cost in counting backwards and subtracting serial sevens. This association remained significant after adjusting for MoCA score, a measure of global cognition (Table 3).

**Discussion**

Our findings reveal that, in a well-characterized cohort of older adults with MCI, a lower volume of the left entorhinal cortex is associated with slow dual-task velocity and higher dual-task gait cost, but not with single-task slow gait. This brain region may be key in explaining the link between dual-task gait performance and progression to dementia.

Our results are aligned with the recent finding that increased dual-task gait cost is associated with an increased risk of progression to dementia in older adults with MCI (12). Recently, lower hippocampal volume was associated with slowing gait in a large-population cohort study (22). This may suggest that dual-task gait performance could be an early motor manifestation of entorhinal cortical atrophy, and when the hippocampal volume is involved, slowing of single gait may become subsequently evident.

Although it is well recognized that both the entorhinal cortex and the hippocampus are affected early in AD pathology (13,14) and that MRI shows abnormalities at very early stages (15–17), a growing body of literature indicates that entorhinal cortex atrophy precedes hippocampal atrophy in pathological aging (36–39). Furthermore, in older adults with MCI, the entorhinal cortex had greater volume loss than the hippocampus (38), and both the volume and rate of atrophy in the entorhinal cortex, but not the hippocampus, were associated with progression to AD (39). Because volume reduction in the entorhinal cortex may be an early sign of AD pathology, our findings may provide an anatomical substrate to the recent finding that lower dual-task gait performance is associated with progression from MCI to dementia (12). Although the association of lower volume in the entorhinal cortex with higher dual-task gait cost was evident in MCI cases, caution should be exercised when drawing conclusions because this association might only be observed in those with a higher risk or progression to dementia.

We found that the relationship between lower volume in the entorhinal cortex and higher dual-task gait cost was observed in the left hemisphere only. It has been suggested that the left entorhinal cortex plays an important role in memory performance (40,41), which would support the concept that other cognitive domains beyond executive function (eg, memory) are involved in controlling

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**Table 2. Correlation Coefficient Matrix of Gait Variables and Regional Brain Volumes**

<table>
<thead>
<tr>
<th></th>
<th>Left Hemisphere</th>
<th></th>
<th>Right Hemisphere</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HC</td>
<td>PHG</td>
<td>EC</td>
<td>MC</td>
</tr>
<tr>
<td>Gait velocitya</td>
<td>.05</td>
<td>.36</td>
<td>.43</td>
<td>.10</td>
</tr>
<tr>
<td>Single gait</td>
<td>.14</td>
<td>.32</td>
<td>.56*</td>
<td>-.02</td>
</tr>
<tr>
<td>Counting gait</td>
<td>.09</td>
<td>.29</td>
<td>.48*</td>
<td>-.08</td>
</tr>
<tr>
<td>Serial sevens gait</td>
<td>.13</td>
<td>.23</td>
<td>.55*</td>
<td>-.10</td>
</tr>
<tr>
<td>Naming animals gait</td>
<td>.21</td>
<td>.19</td>
<td>.49*</td>
<td>.21</td>
</tr>
<tr>
<td>Dual-task costb</td>
<td>-.16</td>
<td>-.28</td>
<td>-.46*</td>
<td>.20</td>
</tr>
<tr>
<td>Counting gait</td>
<td>.21</td>
<td>.01</td>
<td>-.30</td>
<td>.21</td>
</tr>
<tr>
<td>Serial sevens gait</td>
<td>.01</td>
<td>-.01</td>
<td>-.30</td>
<td>.21</td>
</tr>
</tbody>
</table>

Note: EC = entorhinal cortex; HC = hippocampus; LFC = lateral frontal cortex; MC = motor cortex; PHG = parahippocampal gyrus. Correlation coefficients were adjusted for intracranial volume.

aPartial correlation analysis. bPartial rank correlation analysis.

*p < .005.

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**Figure 2.** Scatter graphs of association between dual-task cost in counting backwards and subtracting serial sevens conditions and volume in the left entorhinal cortex.
intracranial volume, years of education, and Montreal Cognitive Assessment.

Serial sevens velocity

in delayed recall among MCI participants (10). Therefore, lower

a close association between dual-task gait cost and performance

and maintaining a safe gait in MCI. A previous study has shown a
close association between dual-task gait cost and performance in
delayed recall among MCI participants (10). Therefore, lower

volume of the entorhinal cortex might affect dual-tasking ability

through memory decline.

In the present study, single-task gait velocity did not show any
association with regional brain volumes. This is inconsistent with
previous findings that slower gait velocity was associated with
reduced gray matter volume in the prefrontal regions and medial
temporal area (18,19,22). A possible explanation is that dual-task

may uncover valuable subtleties regarding the role of cognitive
control in a participant's gait, which gait velocity alone may not be
able to capture, particularly in high-functioning older adults, as in
our sample (12).

Although our study provides novel evidence of a potential mecha-
nism underlying the relationship between the aging brain and dual-
task gait performance among older adults with MCI, some studies
have raised the possibility that brain areas other than the entorhinal
cortex are involved in controlling dual tasking in MCI populations.
A study using magnetic resonance spectroscopy and MRI showed
that the metabolite ratios and volume of the primary motor cor-
tex were associated with gait performance during the dual-task con-
dition (26). More recently, it was shown that in older adults with
MCI a slower gait velocity during the dual-task condition was associ-
ated with a decreased volume in several cerebral regions, includ-
ing the medial frontal gyrus, superior frontal gyrus, and cingulate
(42). Future studies are needed to examine the interactive effects
of atrophy in each brain region on the progression to dementia among
older adults with MCI, focusing on its association with dual-task
gait performance.

The strength of this study is that it is the first to assess the rela-
tionship between dual-task gait velocity and cost in the medi-
tal temporal areas, which are vulnerable regions in AD pathology, in
a well-characterized MCI cohort with detailed evaluations. However,
there are limitations that need to be considered when interpreting
the results. First, the cross-sectional study design precludes our abil-
ity to assess causality. Second, we included a relatively small
number of participants, which may affect the statistical power of
the associations between additional brain regions and dual-task gait
performance. Furthermore, our sample was composed of relatively
well-educated and high-functioning older adults (i.e., mean gait
velocity > 100 cm/s), which limits the generalizability of our results.
Third, because the present study investigated the association of
dual-task cost with regional brain volume, specifically, medial
temporal area, we did not perform whole-brain analysis including
ventricular volume (43,44), which raises the possibility of type 1

error. Fourth, having the cost in the cognitive component of dual

tasking in our cohort could have improved our findings by consid-
ering cognitive errors on dual tasking. Finally, cross-

validation of our findings may be required from other MCI cohorts.

Conclusion

The present study reveals that a lower volume of the entorhinal
cortex is associated with poor dual-task gait performance in older
adults with MCI. Cognitive and motor dysfunction in older adults
with MCI may reflect a shared pathogenic mechanism at the brain
level, and dual-task-related gait changes may be a surrogate motor
marker for AD pathology, such as entorhinal atrophy. Further lon-
gitudinal studies are needed to confirm our results in a larger
population.

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Conflict of Interest

None reported.
References


