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## Original Article

## Relationship between Cortical Thickness and Neuropsychological Performance in Normal Older Adults and Those with Mild Cognitive Impairment

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ABSTRACT: Mild cognitive impairment (MCI) has been extensively investigated in recent decades to identify groups with a high risk of dementia and to establish effective prevention methods during this period. Neuropsychological performance and cortical thickness are two important biomarkers used to predict progression from MCI to dementia. This study compares the cortical thickness and neuropsychological performance in people with MCI and cognitively healthy older adults. We further focus on the relationship between cortical thickness and neuropsychological performance in these two groups. Forty-nine participants with MCI and 40 cognitively healthy older adults were recruited. Cortical thickness was analysed with semiautomatic software, Freesurfer. The analysis reveals that the cortical thickness in the left caudal anterior cingulate (p=0.041), lateral occipital (p=0.009) and right superior temporal (p=0.047) areas were significantly thinner in the MCI group after adjustment for age and education. Almost all neuropsychological test results (with the exception of forward digit span) were significantly correlated to cortical thickness in the MCI group after adjustment for age, gender and education. In contrast, only the score on the Category Verbal Fluency Test and the forward digit span were found to have significant inverse correlations to cortical thickness in the control group of cognitively healthy older adults. The study results suggest that cortical thinning in the temporal region reflects the global change in cognition in subjects with MCI and may be useful to predict progression of MCI to Alzheimer's disease. The different pattern in the correlation of cortical thickness to the neuropsychological performance of patients with MCI from the healthy control subjects may be explained by the hypothesis of MCI as a disconnection syndrome.

*Key words:* cortical thickness, dementia, mild cognitive impairment, neuropsychological performance, magnetic resonance imaging

The significant growth in the population with dementia has been highlighted as a public health priority [1]. A wide range of cognitive impairment is the core symptom of dementia and determines the loss of independent functioning. Mild cognitive impairment (MCI) is a transitional state between normal ageing and dementia [2]. MCI has been extensively investigated in recent decades to identify those with a high risk of dementia and to

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establish effective prevention methods during this period. Neuropsychological performance and cortical thickness are two important biomarkers used to predict progression from MCI to dementia.

Sub-normative neuropsychological performance is one of the core diagnostic criteria for MCI. A wide range of cognitive impairment, including memory, attention and executive functions, can be found in patients with MCI. In addition to its diagnostic value, neuropsychological assessment also provides a possible means of differentiating high-risk groups for different types of dementia [3].

Along with the rapid development of neuroimaging techniques, the use of cortical thickness as measured on T1-weighted magnetic resonance imaging (MRI) as a biomarker to predict or facilitate early diagnosis of dementia has become a research direction of great interest. Compared to voxel-based morphology (VBM), the measurement of cortical thickness allows more precise measurement in deep sulci and analysis of the morphology as a cortical sheet [4]. Convergent findings strongly suggest a significant difference in cortical thickness amongst normal control patients, those with MCI and those with dementia [5-7]. Furthermore, a longitudinal study of 382 participants who were followed up for 24 months suggested that cortical thickness was sensitive for the early diagnosis of Alzheimer's disease [8]. Another study reported that a decrease in cortical thickness could be detected in cognitively normal individuals several years before the onset of clinical symptoms [9].

Cortical thickness was suggested to have a close relationship with neuropsychological performance [10]. Despite the consistent evidence in support of this hypothesis, large variations were found across studies in the correlation of cortical thickness to neuropsychological performance amongst normal older adults and those with MCI and AD. Verbal memory performance was found to be associated with the medial temporal cortical thickness in normal subjects [11]. In subjects with MCI, the thickness of the entorhinal and praecuneus cortices predicted learning, whereas the posterior cingulate cortical thickness predicted learning in subjects with AD [12]. Another study suggested that MCI entails a specific cortical thinning relationship with high-level executive outcomes that is qualitatively different from that observed in healthy older adults [13]. This variation in the correlational patterns may shed light on the underlying differences in the cognitive processes and compensatory mechanisms between people with MCI and normal older adults. There is a paucity of research into differences between people with MCI and healthy subjects in the relationship between neuropsychological performance and cortical thickness. Therefore, we conducted this study to compare the cortical thickness and neuropsychological performance between subjects with MCI and healthy older adults. The relationship between the cortical thickness and neuropsychological performance in these two groups was also examined. We hypothesised that subjects with MCI would have thinner cortices and would display worse neuropsychological performance than healthy older adults. The correlation between the brain cortical thickness and a specific neuropsychological performance may have different patterns in these two groups.

#### MATERIALS AND METHODS

### Subjects

Forty-nine patients with MCI and 40 cognitively healthy elderly control subjects (healthy controls; HC) were recruited. All of the participants were recruited from local elderly community centres. The study was approved by the Clinical Research Ethics Committee of The Chinese University of Hong Kong (NTEC-CUHK ethics committee). Written informed consent was obtained from all of the participants.

All of the participants underwent a battery of neuropsychological tests to evaluate their cognitive functions.

The Cantonese version of the Mini-Mental State Examination (CMMSE) [14, 15] was used to evaluate general cognitive function. The Clinical Dementia Rating (CDR) [16] scale was used to measure the severity of dementia. The Chinese version of the Alzheimer's Disease Assessment Scale-Cognitive Subscale (ADAS-Cog) [17, 18] was used to assess the global cognitive deficit in patients with MCI. In addition, the forward and backward digit span tests from the Wechsler Adult Intelligence Scale [19] were used to assess the function of short-term memory and working memory, respectively. The Category Verbal Fluency Test (CVFT) [20, 21] was used to examine executive and semantic memory functions. The diagnosis of MCI was made by expert neurologists based on the Mayo Clinic Criteria [2], which includes (1) subjective memory complaints, (2) objective memory impairment (i.e., delayed recall scores of at least 1.5 standard deviations below age- and educationmatched persons with a CDR of 0), (3) intact daily life activities, (4) a CDR score of 0.5 and (5) no clinical dementia (CMMSE score > 22 for older adults with more than 2 years of education, CMMSE score > 20 for older adults with less than 2 years of education and CMMSE score > 19 for older adults with no education [22]. Participants with profound sensory deficits or psychiatric

(i.e., dependence on alcohol or other substances) and/or neurological disorders other than dementia (i.e., head trauma, multiple sclerosis and Parkinson's disease) were excluded.

### MRI acquisition

The MRI images were acquired using a 3 Tesla Philips MRI scanner (Achieva TX, Philips Medical Systems, Best, the Netherlands) with an eight-channel SENSE head coil. A 3D high-resolution T1-weighted anatomical image was obtained for each participant (repetition time [TR] = 7.4 ms; echo time [TE] = 3.4 ms; flip angle =  $8^{\circ}$ ; voxel size =  $1.04 \times 1.04 \times 0.6$  mm<sup>3</sup>).

### Cortical thickness analysis

The image data were exported from the MRI scanner to a personal computer for morphometric analysis. Before analysis, all images were checked for severe head motion. Semi-automatic software, the FreeSurfer version 5.3 software package (<a href="http://surfer.nmr.mgh.harvard.edu">http://surfer.nmr.mgh.harvard.edu</a>), was used to obtain estimates of cortical thickness, which was measured by reconstructing representations of the grey/white matter boundary and the cortical surface and then calculating the distance between those surfaces at numerous points (vertices) across the cortical mantle [23,

24]. Failures in FreeSurfer's initial Talairach alignments were identified by visual inspection of all images and were rectified before reconstruction of the cortical surfaces. Topological defects in the automatically determined grey/white matter boundary were manually corrected. The cortical thickness values of 68 structures based on the Desikan-Killiany atlas were extracted from FreeSurfer [25]. All analyses were performed without knowledge of the subjects' identity.

#### Statistical analysis

Linear regression adjusted for age and education was used for statistical analyses of the mean cortical thickness of region of interests (ROIs) between the subjects in the MCI and normal control groups, and p values of less than 0.05 were considered to indicate statistical significance. Partial correlations between neuropsychological scores and mean cortical thickness, adjusted for age, sex and years of education, were calculated for both MCI and control groups. Bonferroni correction was applied to correct for multiple comparisons, and p values of less than 0.01 were considered to indicate statistical significance after correction.

Table 1. Participant demographics and neuropsychological performance.

	Healthy Controls (n=40) Mean (SD)	MCI (n=49) Mean (SD)	p-value
Age	69.45 (4.56)	75.92 (5.39)	< 0.001
Gender (Male: Female)	15:25	26:23	0.143
Education (years)	8.00 (4.00)	4.13 (4.04)	< 0.001
CMMSE	27.68 (2.51)	24.94 (2.85)	< 0.001
CDR – sum of boxes	0.16 (0.43)	1.02 (1.04)	< 0.001
ADAS-Cog	6.46 (2.57)	13.59 (3.61)	< 0.001
Delayed recall	6.58 (1.47)	2.29 (1.46)	< 0.001
CVFT	40.10 (7.58)	31.27 (8.03)	< 0.001
Digit span test (forward)	7.50 (1.36)	6.80 (1.44)	0.021
Digit span test(backward)	3.93 (1.65)	2.59 (1.39)	< 0.01

ADAS-Cog - Chinese version of the Alzheimer's Disease Assessment Scale-Cognitive Subscale; CDR - Clinical Dementia Rating; CMMSE - Cantonese version of the Mini-Mental State Examination; CVFT - Category Verbal Fluency Test

#### **RESULTS**

#### Demographic and baseline data

Table 1 shows significant differences in age and education between the MCI group and the HC group. Compared with those with MCI, the participants in the HC group were younger (mean [SD], 69.45 [4.56] vs. 75.92 [5.39]) and had more years of education (mean [SD], 8.00 [4.00] vs. 4.13 [4.04]). No significant difference was found in the gender ratio. The participants with MCI had significantly lower scores on the CMMSE, CDR sum of boxes, ADAS-Cog, CVFT and forward and backward digit span tests than the subjects in the HC group (p<0.05).

The mean CMMSE score in the MCI group was 24.94, and that in the HC group was 27.68.

## Difference in cortical thickness between MCI and HC groups

The mean cortical thicknesses of all areas in the brain are shown in Table 2 for the MCI group and the HC group.

Analysis reveals significantly less cortical thickness in the left caudal anterior cingulate (p=0.041), left lateral occipital (p=0.009) and right superior temporal (p=0.047) areas in the MCI group after adjustment for age and education.

**Table 2.** Cortical thickness in healthy control and mild cognitive impairment (mean +/- S.D., mm, adjusted for age and education).

	Healthy	Control	N	MCI			
Brain region	Left	Right	Left	Right			
Caudal anterior cingulate gyrus	2.689 (0.315) *	2.599 (0.296)	2.502 (0.378)*	2.512 (0.290)			
Caudal middle frontal gyrus	2.258 (0.168)	2.262 (0.148)	2.218 (0.131)	2.243 (0.145)			
Cuneus	1.618 (0.125)	1.619 (0.118)	1.612 (0.125)	1.606 (0.117)			
Entorthinal area	3.403 (0.392)	3.605 (0.487)	3.288 (0.340)	3.522 (0.413)			
Fusiform gyrus	2.639 (0.148)	2.603 (0.156)	2.577 (0.158)	2.554 (0.188)			
Inferior parietal lobe	2.164 (0.123)	2.115 (0.113)	2.142 (0.135)	2.122 (0.148)			
Inferior temporal gyrus	2.695 (0.161)	2.681 (0.154)	2.613 (0.158)	2.636 (0.184)			
Isthmus cingulate gyrus	2.416 (0.187)	2.302 (0.225)	2.267 (0.229)	2.195 (0.206)			
Lateral occipital gyrus	1.902 (0.130)*	1.879 (0.126)	1.899 (0.152)*	1.874 (0.147)			
Lateral orbitofrontal gyrus	2.522 (0.140)	2.469 (0.153)	2.510 (0.164)	2.430 (0.166)			
Lingual gyrus	1.787 (0.118)	1.810 (0.087)	1.782 (0.144)	1.779 (0.167)			
Medial orbitofrontal gyrus	2.283 (0.170)	2.369 (0.164)	2.289 (0.181)	2.612 (0.165)			
Middle temporal gyrus	2.670 (0.172)	2.746 (0.139)	2.660 (0.142)	2.715 (0.169)			
Parahippocampal gyrus	2.535 (0.230)	2.557 (0.256)	2.378 (0.303)	2.489 (0.264)			
Paracentral gyrus	2.271 (0.179)	2.270 (0.158)	2.223 (0.179)	2.222 (0.158)			
Pars opercularis	2.357 (0.173)	2.366 (0.135)	2.351 (0.120)	2.352 (0.142)			
Pars orbitalis	2.539 (0.217)	2.509 (0.235)	2.471 (0.221)	2.494 (0.247)			
Pars triangularis	2.245 (0.134)	2.279 (0.148)	2.202 (0.134)	2.213 (0.162)			
Periphery calcarine	1.385 (0.878)	1.427 (0.103)	1.414 (0.123)	1.446 (0.128)			
Postcentral gyrus	1.819 (0.132)	1.765 (0.104)	1.779 (0.123)	1.787 (0.118)			
Posterior cingulate gyrus	2.440 (0.221)	2.395 (0.198)	2.345 (0.175)	2.325 (0.177)			
Precentral gyrus	2.364 (0.151)	2.343 (0.124)	2.312 (0.136)	2.284 (0.144)			
Precuneus	2.128 (0.141)	2.064 (0.119)	2.086 (0.161)	2.047 (0.141)			
Rostral anterior cingulate gyrus	2.820 (0.199)	2.882 (0.248)	2.744 (0.223)	2.802 (0.286)			
Rostral middle frontal gyrus	2.110 (0.137)	2.154 (0.120)	2.090 (0.141)	2.139 (0.139)			
Superior frontal gyrus	2.518 (0.146)	2.540 (0.142)	2.475 (0.141)	2.503 (0.137)			
Superior parietal lobe	1.884 (0.135)	1.843 (0.122)	1.863 (0.126)	1.831 (0.121)			
Superior temporal gyrus	2.563 (0.146)	2.596 (0.177)*	2.491 (0.161)	2.574 (0.155)*			
Supramarginal gyrus	2.298 (0.126)	2.229 (0.149)	2.219 (0.141)	2.201 (0.135)			
Frontal pole	2.671 (0.263)	2.634 (0.210)	2.597 (0.256)	2.593 (0.275)			
Temporal pole	3.638 (0.267)	3.759 (0.301)	3.513 (0.283)	3.625 (0.293)			
Transverse temporal gyrus	2.148 (0.252)	2.106 (0.254)	2.070 (0.197)	2.107 (0.203)			
Insula	2.891 (0.157)	2.879 (0.175)	2.861 (0.158)	2.800 (0.165)			

<sup>\*</sup>p<0.05

Table 3. Correlation between neuropsychological performance and cortical thickness in mild cognitive impairment.

Brain region	CMMSE		CDR-Sum of boxes		ADAS-Cog		CVFT		Forward digit span		Backward Digit span	
	Left	Right	Left	Right	Left	Right	Left	Right	Left	Right	Left	Right
Caudal anterior	077	.075	005	.042	139	047	.041	.075	108	213	104	.172
cingulate gyrus												
Caudal middle	202	171	.309	.302	061	.142	108	152	022	184	059	.018
frontal gyrus												
Cuneus	050	.032	062	051	.062	097	.209	.221	151	090	.065	.120
Entorthinal area	.173	.323	262	366	*413	259	.349	.335	.101	.246	301	228
Fusiform gyrus	.213	.239	159	337	137	204	.106	.178	.156	.162	125	016
Inferior parietal	.096	.029	.091	.083	.003	.058	.163	.117	109	110	059	.074
lobe												
Inferior temporal	.191	*.508	023	198	216	369	.350	.262	.023	.214	102	.101
gyrus												
Isthmus cingulate	.336	.201	118	116	277	193	043	032	.115	.134	.120	.246
gyrus												
Lateral occipital	.075	017	133	.005	035	040	.085	016	.007	.092	.085	.225
gyrus	.07.5	.017	.155	.002	.000	.0.0	.002	.010	.007	.0,2	.000	
Lateral	028	040	.202	.003	053	044	.231	.125	.085	038	.176	021
orbitofrontal gyrus	020	040	.202	.003	055	04-4	.231	.123	.003	030	.170	021
Lingual gyrus	.108	.185	111	060	119	226	.172	.209	.048	.125	.165	.191
Medial	.076	.046	023	.033	.030	148	.334	.376	.038	.066	.118	.047
orbitofrontal gyrus	.070	.040	023	.033	.030	140	.554	.570	.036	.000	.116	.047
Middle temporal	.212	.359	.084	182	025	.048	.131	.137	072	083	137	.180
	.212	.339	.064	102	023	.046	.131	.137	072	065	137	.160
gyrus Danahinna a aanna al	215	200	* 412	217	061	102	111	012	004	121	227	100
Parahippocampal	.215	.200	<u>*413</u>	317	061	193	111	012	.004	.131	337	190
gyrus Paragontrol gyrus	144	.043	122	.197	073	044	.005	.048	244	098	.005	.054
Paracentral gyrus	.031	.043	.122 .160	.197	073 031	044 117	.003	.126	244 117	098	.003	187
Pars opercularis												
Pars orbitalis	013	.059	.311	.221	.029	.099	175	050	.277	.245	.315	*.408
Pars triangularis	.045	.051	.058	009	124	155	.170	.159	.013	.138	.227	.302
Pericalcarine	029	251	.048	.072	069	117	.173	.194	.010	.010	.194	.038
Postcentral gyrus	170	188	.131	.107	013	.035	.049	.247	114	146	.042	.081
Posterior cingulate	.039	.040	012	.096	116	.057	.100	036	046	120	067	.181
gyrus							0.40					
Precentral gyrus	044	165	.011	.093	.012	088	040	.003	193	066	090	044
Precuneus	.102	.134	032	.060	151	115	.203	.184	021	106	.032	.022
Rostral anterior	028	070	.284	.239	.024	.214	.016	148	096	090	.087	014
cingulate gyrus												
Rostral middle	267	100	.185	.116	.071	006	.260	<u>*.398</u>	.017	103	.053	.023
frontal gyrus												
Superior frontal	196	190	.391	.255	.032	002	.008	.048	210	098	.007	119
gyrus												
Superior parietal	.002	.029	024	.020	089	028	.151	.188	008	004	.128	.260
lobe												
Superior temporal	.247	.232	142	242	089	.036	.324	.235	.086	.084	.127	.050
gyrus												
Supramarginal	.092	.026	.167	027	174	039	.175	.112	021	.109	.062	.149
gyrus												
Frontal pole	.104	.176	150	089	.042	080	.047	.324	.056	049	.323	005
Temporal pole	.115	.256	215	175	187	209	.356	252	.208	.085	.021	041
Transverse	267	188	.198	.089	.131	.251	.029	170	173	002	.254	.156
temporal gyrus												
Insula	.092	.116	.237	012	162	299	.276	.322	051	.079	.007	078

<sup>\*</sup> p<0.01. ADAS-Cog - Chinese version of the Alzheimer's Disease Assessment Scale–Cognitive Subscale; CDR - Clinical Dementia Rating; CMMSE

## Correlation between cortical thickness and neuropsychological performance in MCI group

Almost all neuropsychological performance, except for the forward digit span, was significantly correlated with the cortical thickness (Table 3). The CMMSE score showed a significant correlation with the right inferior temporal gyrus (r=0.508; p<0.01; Fig. 1). The CDR sum of boxes score showed a significant correlation with the left parahippocampal gyrus (r=-0.413; p<0.01; Fig. 2).

The performance on the ADAS-Cog showed a significant correlation with the left entorhinal area (r=-0.413; p<0.01). The CVFT score showed a significant correlation with the right rostral middle gyrus (r=0.398; p<0.01). Scores on the backward digit span test showed significant correlations with the right pars orbitalis (r=0.408; p<0.01). A thicker cortex in these regions was associated with better performance on the CVFT and on the backward digit span test.

<sup>-</sup> Cantonese version of the Mini-Mental State Examination; CVFT - Category Verbal Fluency Test

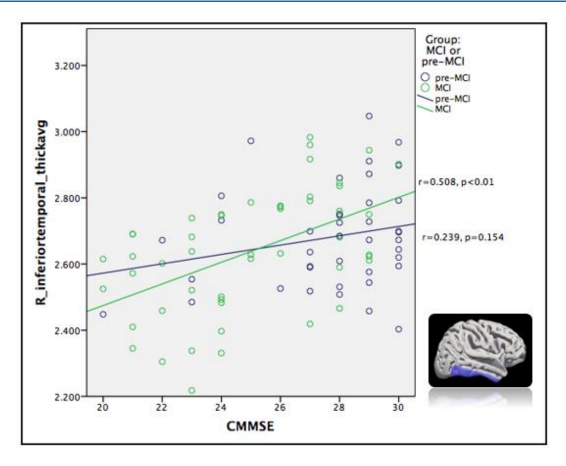


Figure 1. Correlation between right temporal gyrus and Cantonese version of the Mini-Mental State Examination (CMMSE).

## Correlation between cortical thickness and neuropsychological performance in HC group

Only the scores on the CVFT and the forward digit span test were found to have significant correlations with cortical thickness in the HC group (Table 4). The CVFT score showed an inverse correlation with the left middle temporal gyrus (r=-0.445; p<0.01), whilst the forward digit span test score showed a significant inverse correlation with the left pars opercularis (r=-0.496; p<0.01), the left rostral middle frontal gyrus (r=-0.422; p<0.01) and the right orbitofrontal cortex (r=-0.456; p<0.01). A thicker cortex in these regions was associated with poorer performance on the CVFT and on the forward digit span test.

#### DISCUSSION

In this study, we compared the differences in cortical thickness between participants with MCI and those in an HC group. We also examined the association between neuropsychological performance and cortical thickness.

The neuropsychological performance of the MCI group was significantly worse than that of the HC group, which was expected. We found significant thinning in the anterior cingulate and superior temporal regions in participants with MCI compared with those in the HC group. This result is in line with the results of previous studies [26]. It was suggested that cortical thinning begins in the temporal region and spreads to other areas [27]. In addition, the anterior cingulate region was reported in a previous study to be more sensitive comparing to other brain regions to early AD-related changes [6]. Both features were noted in our findings. The cortical thicknesses of these two areas may be useful for early identification of subjects with MCI. In addition to these two areas, the left lateral occipital region was found to be significantly thinner in the MCI group. It was relatively uncommon to note atrophy in the occipital region in subjects with MCI, but studies have nonetheless shown significant increases in the atrophy rate of the occipital region in subjects with AD and MCI [28].

**Table 4.** Correlation between neuropsychological performance and cortical thickness in healthy control.

Brain region         Left         Right           Caudal anterior cingulate gyrus         .010        062           Caudal middle frontal gyrus        066         .116           Cuneus        060         .041           Entorthinal area        252        175           Fusiform gyrus        015        027           Inferior parietal        055         .128           lobe         Inferior temporal gyrus        006         .239           Inferior temporal gyrus        006         .239           gyrus         Lateral occipital         .229         .277           gyrus         Lateral occipital gyrus        079         .222           Medial        191        068           orbitofrontal gyrus        079         .222           Medial        191        068           orbitofrontal gyrus        080        151           gyrus         Parahippocampal        125         .043           gyrus         Paracentral gyrus         .096        086           Pars opercularis        211         .117           Pars orbitalis        228        187           Pars triangularis	Left159 .220 .185 .161 .122 .158 .002	Right177 .164 .025 .162 .131012	.182 .169 .272 .268	Right .257 .106 .095	Left .131 031	<b>Right</b> 166	Left .110	<b>Right</b> 063	<b>Left</b> 076	Right
Caudal anterior cingulate gyrus         .010        062           Caudal middle frontal gyrus        066         .116           Cuneus        060         .041           Entorthinal area        252        175           Fusiform gyrus        015        027           Inferior parietal        055         .128           lobe         Inferior temporal        006         .239           gyrus         Isthmus cingulate        139        137           gyrus         Lateral occipital         .229         .277           gyrus         Lateral occipital         .29         .234           orbitofrontal gyrus         .079         .222           Medial        191        068           orbitofrontal gyrus         .043         .97           Middle temporal        125         .043           gyrus         Paracentral gyrus         .096        086           Pars opercularis        211         .117 </th <th>.220 .185 .161 .122 .158</th> <th>177 .164 .025 .162 .131</th> <th>.182 .169 .272</th> <th>.106 .095</th> <th></th> <th>166</th> <th>.110</th> <th>063</th> <th>- 076</th> <th></th>	.220 .185 .161 .122 .158	177 .164 .025 .162 .131	.182 .169 .272	.106 .095		166	.110	063	- 076	
Caudal middle frontal gyrus        066         .116           Cuneus        060         .041           Entorthinal area        252        175           Fusiform gyrus        015        027           Inferior parietal        055         .128           lobe         Inferior temporal        006         .239           gyrus         Isthmus cingulate        139        137           gyrus         Lateral occipital         .229         .277           gyrus         Lateral occipital         .229         .227           gyrus         Lateral occipital         .229         .227           gyrus         Lateral occipital         .229         .227           gyrus         Lighthus        079         .222           Medial        191        068	.185 .161 .122 .158	.025 .162 .131	.169 .272	.095	031				070	133
Frontal gyrus Cuneus060 .041 Entorthinal area252175 Fusiform gyrus015027 Inferior parietal055 .128 lobe Inferior temporal006 .239 gyrus Isthmus cingulate139137 gyrus Lateral occipital .229 .277 gyrus Lateral278234 orbitofrontal gyrus Lingual gyrus079 .222 Medial191068 orbitofrontal gyrus Middle temporal125 .043 gyrus Parahippocampal180151 gyrus Paracentral gyrus096086 Pars opercularis211 .117 Pars orbitalis228 .187 Pars triangularis200038 Pericalcarine177109 Postcentral gyrus061046 gyrus Precentral gyrus049 .016 Precuneus .059028 Rostral middle276 .054 frontal gyrus Superior frontal .063001 gyrus Superior parietal .092037	.185 .161 .122 .158	.025 .162 .131	.169 .272	.095	031					
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Cuneus        060         .041           Entorthinal area        252        175           Fusiform gyrus        015        027           Inferior parietal        055         .128           lobe        006         .239           Inferior temporal        006         .239           gyrus	.161 .122 .158	.162 .131	.272							
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Solution	.002		.145	.176	024	.006	156	009	.081	.182
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Sthmus cingulate  139  137     Syrus   Lateral occipital   .229   .277     Syrus   Lateral  278  234     Corbitofrontal gyrus  079   .222     Medial  191  068     Orbitofrontal gyrus   Middle temporal gyrus     Parahippocampal  125   .043     Syrus  180  151     Syrus  180  151     Syrus  211   .117     Pars orbitalis  228  187     Pars orbitalis  228  187     Pars orbitalis  228  187     Pars triangularis  200  038     Pericalcarine  177  109     Postcentral gyrus   .144  103     Posterior cingulate  061  046     Syrus  049   .016     Precuneus   .059  028     Rostral anterior  108  117     cingulate gyrus     Rostral middle  276   .054     frontal gyrus   .063  001     Syrus   Superior parietal   .092  037	377									
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Paracentral gyrus         .096        086           Pars opercularis        211         .117           Pars orbitalis        228        187           Pars triangularis        200        038           Pericalcarine        177        109           Postcentral gyrus         .144        103           Posterior cingulate        061        046           gyrus        049         .016           Precuneus         .059        028           Rostral anterior        108        117           cingulate gyrus        276         .054           Rostral middle frontal gyrus         .063        001           Superior frontal         .063        001           gyrus         Superior parietal         .092        037	.070	007	.109	029	293	201	241	233	040	131
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Precuneus         .059        028           Rostral anterior        108        117           cingulate gyrus         Rostral middle        276         .054           frontal gyrus         Superior frontal         .063        001           gyrus         Superior parietal         .092        037	1.45	220	110	102	021	015	226	102	217	122
Rostral anterior108117 cingulate gyrus Rostral middle276 .054 frontal gyrus Superior frontal .063001 gyrus Superior parietal .092037	.145	.220	.110	.183	031	015	326	183	.217	.132
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gyrus Superior parietal .092037										
Superior parietal .092037	.023	.158	.137	.196	025	117	166	295	.092	.088
lobe	.092	.100	.168	.173	.039	.021	074	173	.267	.271
Superior temporal .136 .213	.093	188	.202	.177	.008	029	142	.205	.011	.126
gyrus										
Supramarginal .093020	.134	105	.187	.045	096	103	084	092	.056	.038
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Frontal pole051 .030	051	074	.037	174	238	.074	132	086	.231	.259
Temporal pole007 .131		069	.080	021	198	226	.025	.076	.105	.101
Transverse033 .232	037	289	.131	.051	.022	.235	118	.214	.026	.260
temporal gyrus	037 .051									
Insula .057078		.206	060	.132	220	274	155	229	008	083

\* p<0.0

ADAS-Cog - Chinese version of the Alzheimer's Disease Assessment Scale-Cognitive Subscale; CDR - Clinical Dementia Rating; CMMSE - Cantonese version of the Mini-Mental State Examination; CVFT - Category Verbal Fluency Test

# Correlation between cortical thickness and neuropsychological performance in subjects with MCI

Global cognition as measured by the CMMSE and the CDR sum of boxes showed a moderate correlation with the temporal area in participants with MCI; temporal atrophy is a hallmark of early AD-related changes.

Therefore, our finding supports the notion that cortical thinning in this region is directly linked to a decline in global cognition. This may further support the use of the cortical thickness of the temporal area to predict the progression of MCI to AD.

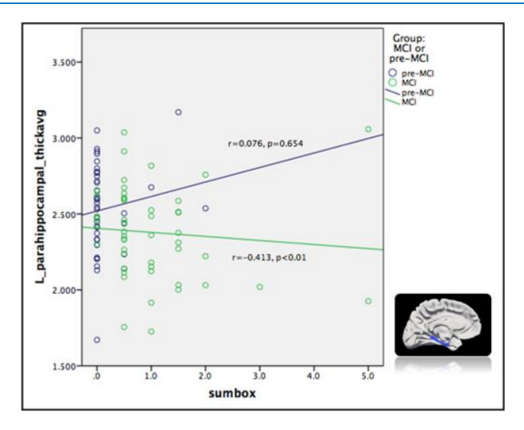


Figure 2. Correlation between left parahippocampal gyrus and Clinical Dementia Rating (CDR)-sum of boxes.

### Difference in correlational patterns

The participants with MCI showed significant correlations between the cortical thickness in various brain areas and each of the neuropsychological performance measures, with the exception of the forward digit span test, but normal older adults showed significant correlations between cortical thickness and two neuropsychological measures only. No global cognition scores such as those on the CMMSE or the CDR sum of boxes were found to have a significant correlation with the cortical thickness in the HC group. One possible explanation for this finding is the ceiling effect of neuropsychological measures in the HC group. However, it could not explain the lack of correlation in tests such as the ADAS-Cog, CVFT and the forward digit span test, in which no prominent ceiling effects were noted. Another postulation is that the participants in the HC group had better connectivity across the whole brain and, therefore, a better compensatory mechanism. When one brain area appeared to be dysfunctional due to the loss of grey matter, other brain areas could compensate so neuropsychological performance and global cognition are relatively maintained. In participants with MCI, due to the

lower degree of connectivity across the whole brain, neuropsychological performance and global cognition directly reflected the severity of cortical thinning without compensation by other brain areas.

The second possible explanation is supported by recent research findings suggesting that MCI and AD represent a disconnection syndrome and that the cognitive impairment results from a decrease in the effectiveness of whole-brain connectivity [29, 30]. A growing body of evidence shows an alteration of functional connectivity in patients with MCI and AD, compared with health control subjects [31, 32]. The connectivity is usually increased in the local area or lobe but significantly decreased across different lobes of the brain [33]. In addition to the functional connectivity, alteration of the structural connectivity, as measured by white matter integrity, has also been reported in patients with prodromal AD [34]. Such Weakening of both functional and structural connectivity may affect the compensatory mechanism. The efficiency of the brain's function as a single unit may decrease. Cognitive functions compartmentally dependent upon one or two areas and are more susceptible to degeneration and loss of neuronal cells. Further study that involves concomitant structural

and functional connectivity investigation is needed to verify that the difference in the relationship between regional cortical thickness and neuropsychological performance between healthy and MCI subjects is due to changes in connectivity.

In our study, scores on the CVFT and the digit backward span test showed a positive correlation with cortical thickness in the MCI group. This means that a decrease in cortical thickness is associated with poorer performance on neuropsychological tests, which is compatible with our previous hypothesis. neuropsychological performance may be more dependent upon the integrity of grey matter in specific brain regions in subjects with MCI due to the impairment of wholebrain connectivity. However, the HC group members had the opposite result: the CVFT and the forward digit span test scores showed a negative correlation with cortical thickness, which means that an increase in cortical thickness is associated with poorer performance on neuropsychological tests. The Previous study also showed that the positive correlation between brain volume and cognition was not found in healthy subjects [35]. One of the possibilities is that the neuropsychological tests were not sensitive enough to reflect the changes in the preclinical phase. The healthy subjects may have AD pathology without symptoms. The previous study found neuronal hypertrophy in the hippocampus and anterior cingulate gyrus neurons among asymptomatic AD patients compared with MCI and control, which may be due to compensation at the local level [36]. Such local compensation may increase cortical thickness but have limited effect on the neuropsychological performance, causing the negative correlation between cortical thickness and neuropsychological performance. However, we could not confirm this explanation in the current study without measurement of AD pathology in our subjects.

Most cognitive training targets deficits in individual cognitive domains. For example, if someone was noted to have a memory problem, the most direct treatment would be to train the memory domain only. However, the effectiveness of this kind of training is in doubt [37]. The effects of training are often short-lived, and the improvement does not translate to daily functions. This phenomenon may be explained by the theory of the disconnection syndrome. The impairment of cognition is due to the connectivity problem rather than solely due to the loss of function of the individual brain areas responsible for that cognitive function. If this is really the case, the aim of cognitive training should be to enhance brain connectivity instead of training up individual cognitive domains. Such connectivity training may have longer and better effects and could likely be generalised

to improvement in daily functioning. Further study is needed to demonstrate this conceptual idea.

## Limitations of study

There were a few limitations of this study. First, the sample size was relatively small and may result in underpower of the current study to detect the difference between the groups. Besides, the pattern difference in correlation between neuropsychological performance and the cortical thickness between MCI and HC groups was mainly descriptive instead of the direct statistical result in the current study. Further study with larger sample size would be needed in order to perform the direct statistical test for comparing the correlation between two groups because a significant amount of multiple comparisons would be involved. Another limitation of our study is the significant difference in education level and age between the MCI group and the HC group; the participants in the MCI group were older and had lower education levels. Comparison of the two groups and correlational analysis were performed with education and age as co-variates to minimise the effect of a baseline difference between the two groups. At last, we had not done the familywise correction for the cortical thickness comparison, which may increase the chance of the false positive result in the current study.

### Conclusions

Our findings suggest that the MCI group had significant thinning over the right temporal, left anterior cingulate and left lateral occipital regions compared with the HC group. Cortical thinning in the temporal region was associated with the global cognition change in participants with MCI and may be useful to predict the progression of MCI to AD. The different pattern between the MCI group and the HC group in the correlation of cortical thickness to neuropsychological performance may be explained by the hypothesis of MCI as a disconnection syndrome. Further imaging studies such as resting state and diffusion tensor imaging are warranted to investigate the alteration in functional and structural connectivity in subjects with MCI. Treatment for cognitive impairment should be directed to the enhancement of brain connectivity in view of the role that a disconnection problem plays in cognitive decline.

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### References

- [1] Prince MJ, Wimo A, Guerchet MM, Ali GC, Wu YT, Prina M. World Alzheimer Report 2015 The Global Impact of Dementia: An analysis of prevalence, incidence, cost and trends. London: Alzheimer's Disease International; 2015.
- [2] Petersen RC, Stevens JC, Ganguli M, Tangalos EG, Cummings JL, DeKosky ST (2001). Practice parameter: early detection of dementia: mild cognitive impairment (an evidence-based review). Report of the Quality Standards Subcommittee of the American Academy of Neurology. Neurology, 56:1133-1142.
- [3] Schindler SE, Jasielec MS, Weng H, Hassenstab JJ, Grober E, McCue LM, et al. (2017). Neuropsychological measures that detect early impairment and decline in preclinical Alzheimer disease. Neurobiol Aging, 56:25-32.
- [4] Lerch JP, Evans AC (2005). Cortical thickness analysis examined through power analysis and a population simulation. Neuroimage, 24:163-173.
- [5] Ries ML, Carlsson CM, Rowley HA, Sager MA, Gleason CE, Asthana S, et al. (2008). Magnetic resonance imaging characterization of brain structure and function in mild cognitive impairment: a review. J Am Geriatr Soc, 56:920-934.
- [6] Zhao H, Li X, Wu W, Li Z, Qian L, Li S, et al. (2015). Atrophic Patterns of the Frontal-Subcortical Circuits in Patients with Mild Cognitive Impairment and Alzheimer's Disease. PLoS One, 10:e0130017.
- [7] Li C, Wang J, Gui L, Zheng J, Liu C, Du H (2011). Alterations of whole-brain cortical area and thickness in mild cognitive impairment and Alzheimer's disease. J Alzheimers Dis, 27:281-290.
- [8] Querbes O, Aubry F, Pariente J, Lotterie JA, Demonet JF, Duret V, et al. (2009). Early diagnosis of Alzheimer's disease using cortical thickness: impact of cognitive reserve. Brain, 132:2036-2047.
- [9] Pettigrew C, Soldan A, Zhu Y, Wang MC, Moghekar A, Brown T, et al. (2016). Cortical thickness in relation to clinical symptom onset in preclinical AD. Neuroimage Clin, 12:116-122.
- [10] Lezak MD. *Neuropsychological assessment*. USA: Oxford University Press; 2004.
- [11] Dickerson BC, Fenstermacher E, Salat DH, Wolk DA, Maguire RP, Desikan R, et al. (2008). Detection of cortical thickness correlates of cognitive performance: Reliability across MRI scan sessions, scanners, and field strengths. Neuroimage, 39:10-18.
- [12] Walhovd KB, Fjell AM, Dale AM, McEvoy LK, Brewer J, Karow DS, et al. (2010). Multi-modal imaging predicts memory performance in normal aging and cognitive decline. Neurobiol Aging, 31:1107-1121.
- [13] Sanchez-Benavides G, Gomez-Anson B, Quintana M,

- Vives Y, Manero RM, Sainz A, et al. (2010). Problemsolving abilities and frontal lobe cortical thickness in healthy aging and mild cognitive impairment. J Int Neuropsychol Soc, 16:836-845.
- [14] Chiu HF, Lee H, Chung W, Kwong P (1994). Reliability and validity of the Cantonese version of mini-mental state examination-a preliminary study. Hong Kong Journal of Psychiatry, 4:25-28.
- [15] Folstein MF, Folstein SE, McHugh PR (1975). "Minimental state". A practical method for grading the cognitive state of patients for the clinician. J Psychiatr Res, 12:189-198.
- [16] Morris JC (1993). The Clinical Dementia Rating (CDR): current version and scoring rules. Neurology, 43:2412-2414.
- [17] Chu LW, Chiu KC, Hui SL, Yu GK, Tsui WJ, Lee PW (2000). The reliability and validity of the Alzheimer's Disease Assessment Scale Cognitive Subscale (ADAS-Cog) among the elderly Chinese in Hong Kong. Ann Acad Med Singapore, 29:474-485.
- [18] Rosen WG, Mohs RC, Davis KL (1984). A new rating scale for Alzheimer's disease. Am J Psychiatry, 141:1356-1364.
- [19] Wechsler D. Wechsler abbreviated intelligence scale. San Antonio: The Psychological Corporation; 1999.
- [20] Chiu HF, Chan CK, Lam LC, Ng KO, Li SW, Wong M, et al. (1997). The modified Fuld Verbal Fluency Test: a validation study in Hong Kong. J Gerontol B Psychol Sci Soc Sci, 52:P247-250.
- [21] Lam LC, Ho P, Lui VW, Tam CW (2006). Reduced semantic fluency as an additional screening tool for subjects with questionable dementia. Dement Geriatr Cogn Disord, 22:159-164.
- [22] Lam LC, Tam CW, Leung GT, Lui VW, Fung AW, Chiu HF, et al. (2010). Combined clinical and cognitive criteria to identify mild cognitive impairment in a southern Chinese community. Alzheimer Dis Assoc Disord, 24:343-347.
- [23] Dale AM, Sereno MI (1993). Improved Localizadon of Cortical Activity by Combining EEG and MEG with MRI Cortical Surface Reconstruction: A Linear Approach. J Cogn Neurosci, 5:162-176.
- [24] Dale AM, Fischl B, Sereno MI (1999). Cortical surface-based analysis. I. Segmentation and surface reconstruction. Neuroimage, 9:179-194.
- [25] Desikan RS, Segonne F, Fischl B, Quinn BT, Dickerson BC, Blacker D, et al. (2006). An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. Neuroimage, 31:968-980.
- [26] Julkunen V, Niskanen E, Koikkalainen J, Herukka SK, Pihlajamaki M, Hallikainen M, et al. (2010). Differences in cortical thickness in healthy controls, subjects with mild cognitive impairment, and Alzheimer's disease patients: a longitudinal study. J Alzheimers Dis. 21:1141-1151.
- [27] Singh V, Chertkow H, Lerch JP, Evans AC, Dorr AE, Kabani NJ (2006). Spatial patterns of cortical thinning in mild cognitive impairment and Alzheimer's disease.

- Brain, 129:2885-2893.
- [28] Sluimer JD, van der Flier WM, Karas GB, van Schijndel R, Barnes J, Boyes RG, et al. (2009). Accelerating regional atrophy rates in the progression from normal aging to Alzheimer's disease. Eur Radiol, 19:2826-2833.
- [29] Gomez C, Stam CJ, Hornero R, Fernandez A, Maestu F (2009). Disturbed beta band functional connectivity in patients with mild cognitive impairment: an MEG study. IEEE Trans Biomed Eng, 56:1683-1690.
- [30] Koenig T, Prichep L, Dierks T, Hubl D, Wahlund LO, John ER, et al. (2005). Decreased EEG synchronization in Alzheimer's disease and mild cognitive impairment. Neurobiol Aging, 26:165-171.
- [31] Wang J, Zuo X, Dai Z, Xia M, Zhao Z, Zhao X, et al. (2013). Disrupted functional brain connectome in individuals at risk for Alzheimer's disease. Biol Psychiatry, 73:472-481.
- [32] Liang P, Li Z, Deshpande G, Wang Z, Hu X, Li K (2014). Altered causal connectivity of resting state brain networks in amnesic MCI. PLoS One, 9:e88476.
- [33] Das SR, Pluta J, Mancuso L, Kliot D, Orozco S, Dickerson BC, et al. (2013). Increased functional connectivity within medial temporal lobe in mild cognitive impairment. Hippocampus, 23:1-6.
- [34] Lacalle-Aurioles M, Navas-Sanchez FJ, Aleman-Gomez Y, Olazaran J, Guzman-De-Villoria JA, Cruz-Orduna I, et al. (2016). The Disconnection Hypothesis in Alzheimer's Disease Studied Through Multimodal Magnetic Resonance Imaging: Structural, Perfusion, and Diffusion Tensor Imaging. J Alzheimers Dis, 50:1051-1064.
- [35] Chee MW, Chen KH, Zheng H, Chan KP, Isaac V, Sim SK, et al. (2009). Cognitive function and brain structure correlations in healthy elderly East Asians. Neuroimage, 46:257-269.
- [36] Iacono D, O'Brien R, Resnick SM, Zonderman AB, Pletnikova O, Rudow G, et al. (2008). Neuronal hypertrophy in asymptomatic Alzheimer disease. J Neuropathol Exp Neurol, 67:578-589.
- [37] Bahar-Fuchs A, Clare L, Woods B (2013). Cognitive training and cognitive rehabilitation for mild to moderate Alzheimer's disease and vascular dementia. Cochrane Database Syst Rev:Cd003260.