3 Tesla MRI detects accelerated hippocampal volume reduction in postmenopausal women

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**Title:** 3 Tesla MRI Detects Accelerated Hippocampal Volume Reduction in Postmenopausal Women

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**Key Words**: Central nervous system; Estrogen; Hippocampus; Magnetic resonance imaging; Menopause; Voxel-based morphometry

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ABSTRACT

Purpose: To clarify age-related structural changes specific to hippocampal volume by hierarchizing according to age, gender, and menopausal status. Many studies report the neuroprotective effects of estrogen and age-related brain volume changes; however, there are no studies regarding age-related change specific to hippocampal volume in terms of age, gender, and menopausal status.

Materials and Methods: T1-weighted magnetic resonance images were obtained in 412 healthy adults divided into 8 groups according to age and gender, to analyze brain volume change focusing on hippocampal volume.

Results: Voxel-based morphometry (VBM) revealed significantly smaller gray matter volume in the hippocampus bilaterally in females aged in their fifties (51 of 59 females were at menopause) compared with females in their forties (3 of 46 females were at menopause). No significant difference was found, however, between female groups in their fifties vs. sixties, or sixties vs. seventies; or between male groups in their forties vs. fifties, fifties vs. sixties, or sixties vs. seventies. In addition, VBM revealed significant hippocampal volume reduction bilaterally in all post-menopausal women compared with all pre-menopausal women.

Conclusion: The results of the current study suggest that the menopause may be associated with hippocampal volume reduction.
Introduction

Many studies report the neuroprotective effects of estrogen and describe the relation between estrogen therapy and hippocampal volume (1), the association between hippocampal glucose metabolism and cerebrospinal fluid $17\beta$-estradiol concentration in post-menopausal women (2), the lower risk of incident Alzheimer’s disease in post-menopausal women using estrogen (3), and atrophy of hippocampal volume in patients with Alzheimer’s disease (4). However, some studies have suggested otherwise (5–8). For example, Low et al. reported that no differences were observed between the control group and the hormone replacement therapy (HRT) group in terms of total gray matter, white matter, hippocampal or amygdalar volumes, severity or volume of white matter hyperintensities, or in different measures of brain atrophy (5). Furthermore, the neuroprotective role of estrogen remains a controversial issue because analytical results are influenced by the type of menopause (i.e., surgical menopause and natural menopause) and the type of cognitive test (i.e., verbal, visual and other memory) (9).

Numerous studies have investigated age-related structural brain changes using MRI, showing a negative correlation between gray matter volume and age, predominantly in the frontal lobes (10–15). These studies reported that the rate of brain volume decline differed according to the focal region and gender (11,13,16–20), as well as a negative correlation between hippocampal volume and age (12,20,21); however, to the best of our knowledge, no study has reported a correlation between menopause and
hippocampal volume.

The aim of the present study was to investigate hippocampal volume change in normal adults divided into eight groups according to their age and gender, using high spatial resolution T1-weighted images with 3.0 tesla (3.0-T) magnetic resonance scanners and voxel-based morphometry (VBM) (22). In addition, we compared hippocampal volume between the pre-menopausal women and post-menopausal women.
MATERIALS AND METHODS

Subjects

A total of 470 subjects who visited our hospital for a personal health-screening program between October 2006 and March 2007 participated in this study. Subjects with a Mini-Mental State Examination (23) score of 26 or lower were excluded from this study. We were not able to exclude mild cognitive impairment. Age at menopause was obtained from all female subjects at interview. The MR images were inspected by a board-certified radiologist (N.H.), and subjects with the following findings were excluded from the study: brain tumors, infarctions, hemorrhage, and white matter lesions graded higher than grade 2 of Fazekas’s classification (24). A total of 58 subjects were excluded from this study. We divided the remaining 412 subjects into the following groups according to age and sex: females aged in their forties (mean ± standard deviation (SD), 45.1 ± 2.9 years, n = 46), females in their fifties (55.4 ± 2.7 years; n = 59), females in their sixties (64.2 ± 2.6 years; n = 49), females in their seventies (74.1 ± 3.0 years; n = 17), males in their forties (44.9 ± 2.8 years; n = 85), males in their fifties (54.2 ± 3.1 years; n = 80), males in their sixties (64.8 ± 2.7 years; n = 63), and males in their seventies (73.6 ± 3.0 years; n = 13).

Hypertension was defined as maximal blood pressure of 140 mmHg or more, or minimal blood pressure of 90 mmHg or more. Drinking index was defined as amount of alcohol consumed per day (we assumed 33.6 ml ethanol to be equal to 1, i.e. the volume
of ethanol that is approximately equal to one glass of wine) × (number of drinking days per week). Smoking index was equal to the Brinkmann index and was expressed as (number of cigarettes smoked per day) × (years of smoking). These data and the subjects’ characteristics are summarized in Table 1. Although some studies report that brain atrophy is associated with hypertension (25–28), alcoholism (29,30), and blood glucose level (31–33), subjects were not excluded on the basis of clinical information because we did not wish to generate a supernormal group of subjects. Analysis of variance (ANOVA) with the Bonferroni–Dunn method as a post hoc test was used to compare drinking index, smoking index, and blood glucose level among the groups. The Kruskal–Wallis test was used to compare the number of subjects with hypertension among the age groups. In both tests, statistical significance was set as $P$ value < 0.05.

The protocol was approved by the Ethics Committee of our institution. After the study had been explained to each subject, written informed consent was obtained from all participants.

**MRI Scanning Protocol**

MRI data were obtained using a 3.0-T scanner (Signa Excite HDx, GE Medical Systems, Waukesha, WI). An 8-channel phased-array brain coil was used as the receiver coil. Three-dimensional fast spoiled-gradient recalled acquisition in the steady state (3D-
FSPGR was used to obtain 180 contiguous sagittal T1-weighted images with a slice thickness of 1.0 mm for VBM analysis (TR / TE = 5.3 / 1.8 ms; inversion time = 450 ms; flip angle = 15°; field of view = 24 cm; number of excitations = 0.5; 256 × 256 pixel matrix). The voxel dimensions were 0.9375 × 0.9375 × 1.0 mm. An acceleration factor of R = 2.0 was employed for parallel imaging using the Array Spatial Sensitivity Encoding Technique.

Image Preprocessing for VBM

We used Statistical Parametric Mapping 5 (SPM5) software (Wellcome Department of Imaging Neuroscience Group, London, UK; http://www.fil.ion.ucl.ac.uk/spm) (34). The 3D-FSPGR images in native space were bias-corrected, spatially normalized, and segmented into gray matter, white matter, and cerebrospinal fluid images (22); voxel size of the normalized images was 2 × 2 × 2 mm. The affine regularization space template from the International Consortium for Brain Mapping was changed from the European to the East Asian brain template. During the modulation step, the voxel values of the normalized gray and white matter images were multiplied by a measure of the relative volumes of the warped and unwarped structures derived from the nonlinear step of spatial normalization using the Jacobian determinant. The total gray and white matter volume, and the total intracranial volume (TIV; measured in ml) were computed by
multiplying the voxel value by the voxel volume and summing the results for all voxels. The resulting gray matter images were smoothed with a Gaussian kernel of 6 mm full width at half-maximum.

Statistical Analyses for VBM

The normalized data smoothed with 6-mm isotropic Gaussian kernels were analyzed with SPM5 employing the framework of the general linear model. Statistical significance for gray matter volume between the paired sequential age groups (subjects in their forties vs. fifties, fifties vs. sixties, sixties vs. seventies) of both males and females was tested with TIV as the confounding covariate to control for brain volume differences between subjects. To test hypotheses with respect to regionally specific group effects, the estimates were compared with two linear contrasts (increased or decreased gray matter volume). The significance of each region was estimated by distributional approximations from the theory of random Gaussian fields. A $P$ value of less than 0.05 corrected with family-wise error (FWE) in voxel difference and a cluster size greater than 30 voxels was considered to be statistically significant.

In addition, regional gray matter volume of the group of all pre-menopausal women (mean ± SD, 45.8 ± 3.6 years; n = 51) was compared with that of all post-menopausal women (61.3 ± 6.7 years; n = 120) in a voxel-wise manner to avoid type 1 or type 2 error due to multiple pairs of group comparisons (e.g. forties vs. fifties, fifties vs. sixties,
sixties vs. seventies for each sex). Because we hypothesized that menopause might have
a certain effect on hippocampal volume, we employed small-volume correction for
multiple comparisons (20 × 30 × 26-mm regions at center (± 26, –24, –8)) (35). We
treated age and TIV as confounding covariates, in this analysis. Previous neuroimaging
studies reported that hippocampal volume negatively correlates. We treated age as a
confounding covariate to control for age-related volume changes because subjects with
a wide range of age were enrolled in this analysis. A $P$ value of less than 0.05 corrected
with FWE in voxel difference was considered to be statistically significant.
RESULTS

Subjects

ANOVA revealed no significant differences in drinking index ($P = 0.120$), smoking index ($P = 0.204$), or blood glucose level ($P = 0.286$) among the female groups; or in smoking index ($P = 0.948$) or blood glucose level ($P = 0.064$) among the male groups. There was significant difference in drinking index ($P = 0.014$) among the male groups tested with ANOVA. Significant difference was found in drinking index only between the groups of males in their forties vs. sixties ($P = 0.004$) with a post hoc test. Kruskal–Wallis test revealed no significant difference in the number of subjects with hypertension among the female groups ($P = 0.391$) or the male groups ($P = 0.061$).

Differences in Gray Matter Volume Using VBM

Bilateral hippocampal volumes were significantly smaller for females in their fifties than in their forties (Fig. 1). Montreal Neurological Institute (MNI) coordinates of local maxima were 26, −24, −18 (max $P$ value $< 0.001$, $T$ value $= 6.96$, cluster size $= 208$) for the right hippocampus, and −28, −22, −18 (max $P$ value $< 0.001$, $T$ value $= 6.16$, cluster size $= 161$) for the left hippocampus. There was no significant difference in hippocampal volume between the female groups in fifties vs. sixties, sixties vs. seventies, or the male groups in forties vs. fifties, fifties vs. sixties, or sixties vs. seventies. In addition, regional gray matter volume in the group of all pre-menopausal
women was compared with that of all post-menopausal women in a voxel-wise manner. Significant hippocampal volume reduction was found bilaterally in the post-menopausal group compared with the pre-menopausal group, in good agreement with the results of group comparison between females in their forties vs. fifties. MNI coordinates of local maxima were 26, –26, –18 (max $P$ value = 0.031, $T$ value = 4.51) for the right hippocampus, and –30, –18, –24 (max $P$ value = 0.043, $T$ value = 3.59) for the left hippocampus.
DISCUSSION

In the present study, voxel-wise comparison revealed that gray matter volumes in the bilateral hippocampus were significantly smaller in females in their fifties (51 of 59 females were at menopause) than in their forties (only 3 of the 46 females were at menopause). In addition, a significantly smaller hippocampus was found in the post-menopausal group than in the pre-menopausal group. The significant volume reduction found in females in their fifties might have type 1 or type 2 errors due to multiple pairs of group comparisons; however, the significant volume reduction found in the post-menopausal group did not suffer from type 1 or type 2 errors because no multiple comparisons were made. Furthermore, these results were consistent, and complementary to each other. To the best of our knowledge, no previous studies have compared hippocampal volume change in detail according to age and sex. In our study, it is not clear that post-menopause women show faster age-related effects than the pre-menopause women or that there is a sudden drop in hippocampus volume after menopause. We expect that it be clarified by longitudinal study in future investigation.

Many studies have reported a negative correlation between gray matter volume and age, while others have reported that the rate of decline of brain volume differs according to the focal region and sex (11,16-20). Several studies reported findings consistent with age-related reduction in hippocampal volume; however, these reports did not consider the rate of decline between groups divided according to age.
In the present study, hippocampal volume measurements revealed a significant difference bilaterally between the groups of females in their forties vs. fifties; there was no significant difference between the groups of females in their fifties vs. sixties, or in their sixties vs. seventies. We consider that the reason for this discrepancy is that reduced hippocampal volume over one decade was so small compared to the SD in each group that a statistically significant reduction was not found in group comparisons.

Previous longitudinal studies indicate that the process of atrophy differs among individuals (36,37). In other words, within-group variance in people in their forties and fifties is lower than that in people in their sixties and seventies; i.e., comparison of data between people in their sixties and seventies is more insensitive to change than comparison between data of those in their forties and fifties. We were not able to eliminate this problem; however, significant difference was found in brain volume between the female groups in their forties and fifties but not between the male groups in their forties and fifties. We think that the different results between the sexes could be indirect evidence of the effects of menopause on hippocampal volume. Several studies report various rates of hippocampal volume reduction (12,19,20,38) because the rate of reduction might be affected by varying factors in the studies (age range, male–female ratio, ROI, etc.). Therefore, we must interpret the results of the rate of hippocampal volume reduction quantitatively and with much caution.

Lord et al. studied the relation between estrogen therapy and hippocampal volume in
estrogen therapy users, past users, never users, and men, and suggested a positive association between estrogen and hippocampal volume (1). In a study that measured the 17β-estradiol levels in cerebrospinal fluid (CSF) and regional cerebral (18F) 2-fluoro 2-deoxy-D-glucose uptake in six female post-menopausal patients with probable Alzheimer’s disease (mean age = 70.3 ± 7.7 years, age range = 68–78 years), Schonknecht et al. suggested an association between estrogen and hippocampal glucose metabolism (2). Tang et al. reported that estrogen use in post-menopausal women might delay the onset and decrease the risk of Alzheimer’s disease (3) by the neuroprotective effect of estrogen (9,39). In contrast, Amagai et al. studied 4683 females (age range = 36–89 years) and reported the mean age ± SD at menopause as 48.3 ± 4.8, with 80% of females experiencing menopause between 45 and 54 years of age (40). Furthermore, in the present study, only 3 of 46 females in their forties were at menopause, while 51 of 59 females in their fifties were at menopause. Therefore, we speculate that the reduced estrogen levels in post-menopausal women might accelerate the rate of decline in hippocampal volume through forty and fifty years old.

A major limitation of the current study is the lack of information regarding estrogen levels in the blood and in CSF. Unfortunately, because estrogen level was not measured as part of our hospital’s personal health screening program, we are not able to infer a direct relationship between estrogen level and hippocampal volume. However, previous reports indicate that the prevalence of HRT use in Japan is approximately 2% (41).
Therefore, we consider that HRT would have had very little effect on the present results.

Previous studies report that as well as estrogen level, hypertension (15,25–28), drinking index (29,30), smoking index (42,43), and blood glucose level (31–33) have a certain effect on brain volume. We consider, however, that these additional factors would have had very little effect on the present analysis because we found no significant difference in hypertension, drinking index, smoking index, or blood glucose level among the female groups. Furthermore, significant difference was found in brain volume between the female groups in their forties and fifties but not between the male groups in their forties and fifties. We think that these different results between the sexes could be indirect evidence of menopausal effects on hippocampal volume, and we therefore speculate that the reduction in estrogen levels in post-menopausal women accelerates the rate of hippocampal volume decline in the female group in their fifties compared to that in their forties. However, to confirm our prediction, we consider that future investigation is needed, using information regarding estrogen levels and HRT.

The second limitation of our study was that a significant difference in drinking index was found only between the groups of males in their forties vs. sixties. This result suggests that alcohol consumption affects age-related reduction of hippocampal volume and could cause overestimation of the rate of hippocampal volume reduction in males from a quantitative viewpoint; however, a significant difference in drinking index was not seen in the results of female group differences between the forties and fifties. To
investigate the effect of alcohol intake on brain volume, we tested correlations between gray matter volume and drinking index in a voxel-wise manner, treating age and TIV as confounding covariates among all subjects. The results revealed no significant correlation between hippocampal volume and drinking index. Therefore, a significant difference in drinking index between the groups of males in their forties vs. sixties had a minor effect on our morphometric results.

Although hypertension was more prevalent in the older groups, there was no significant difference in the number of persons with hypertension among the female groups or the male groups. Therefore, we did not exclude subjects on the basis of hypertension or drinking because we thought that these factors would have very little effect on the volumetric findings, and we did not wish to generate a supernormal group of subjects. Because we considered influence from third variables, we tried to exclude the influence of various factors (i.e., hypertension, drinking index, smoking index, and blood glucose level); however, we consider that influence from other variables may not have been completely excluded. Therefore, the results of the present study need to be interpreted with caution because third variables may explain the group differences.

We performed an additional ROI study using linear regression and polynomial correlation analyses. These results showed a negative correlation between gray matter volume and age, but were not able to prove a difference within the critical age window for menopausal status; however, when groups were divided according to age and sex,
we detected accelerated reduction in the group of females in their forties compared with fifties. It is arbitrary to group the subjects by decades of age, but correlational analyses (such as linear, polynomial, and logarithmic regression) may not enable detection of volume changes at specific decades of age. In other words, using this method of analysis, we could detect accelerated volume changes in the hippocampus between women in their forties and fifties, but not in other decades. Furthermore, because we treated age as a confounding covariate in our analysis, age-related volume change could be controlled.

In conclusion, VBM revealed significant hippocampal volume reduction only between groups of females in their forties vs. fifties. A similar reduction was found between the pre-menopausal and post-menopausal groups. The results of the current study suggest that the menopause may be associated with hippocampal volume reduction.
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Age, age at menopause, drinking index, smoking index, and blood glucose level are shown as average ± standard deviation.

Hypertension was defined as maximal blood pressure \( \geq 140 \text{mmHg} \) or minimal blood pressure \( \geq 90 \text{mmHg} \).

Drinking index was defined as \( \text{amount of alcohol consumed per day} \times \text{(number of drinking days per week)} \).

Smoking index was equal to the Brinkmann index and was expressed as \( \text{(number of cigarettes smoked per day) } \times \text{(years of smoking)} \).
Figure 1. Areas of decreased gray matter volume between females in their forties vs. fifties. Regions of gray matter volume difference are superimposed onto the T1 template image. The color bar (bottom right) represents the $T$ score. R and L indicate the right and left sides of the subjects, respectively. Right hippocampus: max $T$ value = 6.96, cluster size = 208, $(x, y, z) = (26, -24, -18)$. Left hippocampus: max $T$ value = 6.16, cluster size = 161, $(x, y, z) = (-28, -22, -18)$. 