Original Article
Structural gray matter change early in male patients with HIV

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Abstract: The purpose of this study is to characterize brain gray matter volumetric changes in HIV seropositive without neurocognitive impairment and seronegative men in Asia. We investigate 36 males with HIV seropositive (mean age 34.5±9.1 years) and 33 age- and gender-matched seronegative controls (mean age 31.4±7.6 years) in Asia. The cognitive competence of 36 males with HIV seropositive has no impaired based on performance in the international HIV dementia scale. High-resolution T1-weighted magnetic resonance imaging is performed on a 3.0 T MR system using a standard 32-channel birdcage head coil. Voxel-based morphometry is used to derive volumetric measurements at the level of the individual voxel (p < 0.001, none corrected for multiple comparisons). Compared to the control group, HIV seropositive male lower gray matter volumes are found in left inferior frontal gyrus triangular part and orbital part, left superior temporal gyrus, right middle frontal gyrus and cingulum; significant increases gray matter volumes can be seen in Periaqueductal gray and gray around lateral ventricle. HIV infection can change the gray matter volume early without cognitive competence impaired and MR can recognize that changes.

Keywords: Morphometry, MRI, gray matter, HIV

Introduction

Despite the widening use of highly active anti-retroviral therapy (HAART), neurocognitive impairment remains common among HIV-infected (HIV+) individuals. It has been established that viral invasion of the brain occurs early in infection [1-3], and leads to cognitive impairment [4]. Brain atrophy, and injury to subcortical regions, have been shown later in the course of infection [5-7].

Although various neuroimaging studies investigated structural brain changes in HIV, recent MR-based studies reported volumetric changes of the whole brain, basal ganglia in HIV patients [8-10]. Various correlations of MR parameters to cognitive decline were shown in these studies. However, it remains unclear if HIV positive patients without cognitive decline would also suffer from brain tissue atrophy.

VBM is a method to identify regional brain volume differences of gray matter, white matter and cerebrospinal fluid (CSF) using voxel-wise statistics in the context of Gaussian random fields [11]. The VBM procedure allows the detection of highly localized differences, consistently observed across the samples, over the entire brain, even in areas where the region of interest analysis would be difficult. The procedure has proved to be a powerful method in detecting regional tissue differences, even in clinical conditions where routine imaging does not show any visible abnormality [12, 13], and provides the opportunity for an unbiased general search of abnormalities in the whole-brain volume [11]. Thus VBM may be useful to examine regional gray matter changes in HIV-infected male.

In this study, we have compared a group of HIV-infected males with gender- and age-matched healthy control group with VBM, using high-resolution T1-weighted images and statistical parametric mapping method.

Materials and methods

In this study we enrolled 36 males patients with HIV seropositive (mean age 34.5±9.1 years, range: 18-50 years; mean current CD4 cell count 206±187 cells/ml, range: 2-673 cells/ml;
mean nadir CD4 cell count 199±189 cells/ml; 33 age- and gender-matched HIV negative healthy controls (mean age 31.4±7.6 years, range 22-58 years) without any neurological limitation. All patients were recruited from You’an Hospital (Beijing) between June 2011 and May 2013. Inclusion criteria were HIV seropositive, informed consent and age older than 18 years. Exclusion criteria were opportunistic systemic or CNS infection, CNS neoplasm, active alcohol or drug abuse, concomitant neurological disease caused by factors other than HIV, major depression or other severe psychiatric disease. All patients and controls underwent a careful neurological examination by experienced neurologists. All patients and healthy subjects gave their written informed consent according to the Declaration of Helsinki prior to study inclusion. The study was approved by the local ethics committee.

Neurocognitive testing was performed with the international HIV dementia scale (IHDS) [14]. The IHDS comprises four tasks that evaluate the domains of memory (recall of four items at 5 min), attention (anti-saccadic errors), psychomotor speed (timed written alphabet), and construction. The maximum score achievable in the IHDS is 16 and a score < 10 predict HIV dementia. At this stage, we enrolled patients with no cognitive impairment (IHDS score ≥ 15).

History of HAART was received by 11 patients mean duration 3.3±3.8 years, range: 0.25-11.5 years.

Table 1. Regions of reduced gray matter volumes in HIV positive patients without cognitive impairment compared with age- and gender-matched healthy controls

<table>
<thead>
<tr>
<th>Cluster kE</th>
<th>T value</th>
<th>Coordinates (x, y, z)</th>
<th>Anatomic location</th>
</tr>
</thead>
<tbody>
<tr>
<td>1608</td>
<td>4.58</td>
<td>5 35 24</td>
<td>Right anterior cingulate gyrus</td>
</tr>
<tr>
<td>1515</td>
<td>4.51</td>
<td>-45 47 3</td>
<td>Left inferior front gyrus triangle</td>
</tr>
<tr>
<td>139</td>
<td>4.08</td>
<td>-32 38 -18</td>
<td>Left inferior front gyrus orbital</td>
</tr>
<tr>
<td>196</td>
<td>4.04</td>
<td>-51 -36 21</td>
<td>Left superior temporal gyrus</td>
</tr>
<tr>
<td>197</td>
<td>4.03</td>
<td>32 36 33</td>
<td>Right middle front gyrus</td>
</tr>
<tr>
<td>317</td>
<td>4.03</td>
<td>39 20 45</td>
<td>Right middle front gyrus</td>
</tr>
<tr>
<td>214</td>
<td>3.73</td>
<td>47 44 6</td>
<td>Right middle front gyrus</td>
</tr>
<tr>
<td>126</td>
<td>3.96</td>
<td>51 -51 -32</td>
<td>Right crura cerebelli</td>
</tr>
</tbody>
</table>

Coordinates (x, y, z) refer to the location within the MNI space and denote the cluster peak location (defined as voxel with the highest T value); kE Cluster size expressed in mm³.

Imaging data were performed on a single magnetic resonance scanner, at a 3.0 T MAGNETOM Tim Trio (Siemens, Erlangen, Germany) using a standard 32-channel birdcage head coil. T1-weighted sagittal whole-brain magnetic resonance imaging 3D datasets were obtained using magnetization prepared rapid acquisition gradient echo (MP-RAGE) sequence (TR/TE/TI=1900 ms/2.52 ms/900 ms, flip angle=9°, field of view=246×256 mm, slice thickness=1 mm; resolution=1×1 mm; slices=176). The same scanner and the same scanning protocol were used for all patients and healthy controls.

We used statistical parametric mapping (SPM8) software (Wellcome (or Welcome?) Department of Cognitive Neurology, University College of London, London, England; http://www.fil.ion.ucl.ac.uk/spm) running under MATLAB version 8.0 (MathWorks Inc., Natick, Massachusetts) to data processing and analysis.

The standard VBM method can be summarized by the following sequence of steps: spatial normalization, segmentation, modulation, and smoothing. Pre-processing involved all voxel signal intensities in the final GM segmented images which multiplied by the Jacobian determinants (Jacobian modulation) derived from the spatial normalization [15]; segmentation into gray matter, white matter and CSF, and spatial smoothing with an isotropic Gaussian kernel of 8 mm full-width at half maximum (FWHM) [15]. After smoothing, each voxel of the image was a locally weighted average of GM or WM density from a region of surrounding voxels, and statistical analyses can be performed [16, 17]. VBM analysis was performed in a cross sectional setting regional gray matter volumes of HIV positive patients without cognitive deficits and healthy controls, which were compared in Two-sample t-test analysis. Level of significance was set to a P value of < 0.001 with a cluster threshold of 100 voxels. To avoid edge effects at the border of gray matter, we excluded all voxels with a gray matter value below 0.2 (maximum value 0.8). Results were
Superimposed on a Montreal Neurologic Institute (MNI) template (Colin Brain).

We compared patient and control groups with age using independent t-tests and chi-square tests; when appropriate, nonparametric analogues (Mann-Whitney tests and Fisher’s exact tests) were conducted. All tests were two-tailed with alpha level=0.05.

**Result**

Initially, 131 HIV-positive patients were enrolled in the study. Before processing and analyzing the images with VBM, in accordance with T1WI and T2WI we excluded a total of 76 cases because of subclinical CNS infection and neoplasm, lacunar infarct, image 160 slices and the images have motion artifact or other various artifacts which did not pass our quality control checks; excluded 8 female, and 11 cognitive impairment patients. All these subjects were excluded from further analysis. And then remaining 36 male patients showed no abnormalities in T1WI and T2WI.

**Table 1** gave an overview of gray matter of brain reduces detectable when comparing controls with HIV patients without cognitive decline. Most severe gray matter reduction was present the bilateral anterior cingulate cortex especially

**Table 2.** Regions of increased gray matter volumes in HIV positive patients without cognitive impairment compared with age- and gender-matched healthy controls

<table>
<thead>
<tr>
<th>Cluster kE</th>
<th>T value</th>
<th>Coordinates (x, y, z)</th>
<th>Anatomic location</th>
</tr>
</thead>
<tbody>
<tr>
<td>158</td>
<td>3.84</td>
<td>-5 -16 -14</td>
<td>Periaqueductal gray matter</td>
</tr>
<tr>
<td>134</td>
<td>4.17</td>
<td>-21 -78 18</td>
<td>Periventricular gray matter</td>
</tr>
<tr>
<td>286</td>
<td>4.36</td>
<td>-23 -12 15</td>
<td>Periventricular gray matter</td>
</tr>
<tr>
<td>341</td>
<td>5.06</td>
<td>-3 3 19</td>
<td>Periventricular gray matter</td>
</tr>
<tr>
<td>277</td>
<td>5.53</td>
<td>-14 -45 19</td>
<td>Periventricular gray matter</td>
</tr>
</tbody>
</table>

Coordinates (x, y, z) refer to the location within the MNI space and denote the cluster peak location (defined as voxel with the highest T value); kE Cluster size expressed in mm$^3$.
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the right side, as well as the left inferior frontal gyrus triangular part and orbital part (Figure 1). Furthermore, atrophy was present within left superior temporal gyrus, Right middle frontal gyrus marked volume reduction comprising three separate clusters (Table 1). In the crura cerebelli gray matter reduction was detected on the right side.

Table 2 showed an overview of structural brain gray matter increased when comparing controls with HIV patients without cognitive decline. Most severe gray matter increase was present around the lateral ventricles (Figure 2). Furthermore, Periaqueductal gray matter was increased (Figure 2).

At the beginning of the study, we did a pretest study, and discovered that gender had a significant effect on gray matter volume change (Figure 3). That was the reason why we excluded the female patients from further analysis.

Discussion

It’s well known that clinical symptoms of central nervous system (CNS) disease are based on neuronal dysfunction or death, rather than HIV directly infected neurons. However, the mechanism of HIV-related brain damage is not fully clear. The current theory is that the CNS infection is continuous and self-supporting, and does not depend on subsequent cycle of the spread of the virus.

Before HAART, HIV Associated Dementia (HAD) mainly detected in advanced HIV infectors. With the widespread use of HAART, prevalence of HAD decreased significantly, the health status of patients also improved. However, HIV-associated neurocognitive disorder (HAND) prevalence close to 40% in HIV-infected persons [18-20], is still a serious impact on the health of patients [21]. Therefore, detecting HAND early has a very important clinical significance.
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Although HAART effectively inhibit viral replication and rebuild the immune system, mild HAND can still occur in each period of HIV infections [22, 23]. A study [24] found significant psychological behavior abnormalities in these patients even more than the movement abnormalities. Cognitive impairment is still the main symptom of HAND patients, and the symptom was ingravescent. As we conclude from our IHDS test to these HIV patients, we found that with the disease developing, cognitive impairment become more obvious.

Conventional MR examination can find exact atrophy of cortex, but when it does not find a confirm brain structural abnormalities early, VBM can discover the change of brain volume. Through analysis of the data, we found that gray matter reduction of HIV patients without cognitive impairment was in bilateral anterior cingulate gray matter, the left inferior frontal gyrus triangular part and orbital part, right middle frontal gyrus, a smaller volume of the left superior temporal gyrus, and the cerebellar crus compared to the control group.

With a review of the literatures, we found that, as studies have shown that cognitive impairment of HAND is associated with early all subcortical and frontal lobe damage [25], in our study, the left frontal lobe (including inferior frontal gyrus triangular part and orbital part) brain tissue decreased significantly, these two areas mainly involves memory, cognitive and emotional decision function. We also found that the volume of the left superior temporal lobe cortex tissue is reduced, which is the same as the recent several studies. One of the research on visual attention task by fMRI-BOLD showed that HIV positive patients compared with control group the left temporal lobe signals continue increasing in 1 year [26], proved that the temporal lobe cortex sustained damage in HIV infection; Another study showed that temporal lobe cortex tissue decreased significantly with impaired cognitive function [27], also prompt the function of temporal lobe cortex associated with cognitive function. But the function of the posterior superior temporal lobe is mainly the Wernicke area associated with impaired auditory language comprehension [28], can further induce the symptom of sensory aphasia for HAND. At the same time, we found several volumes decrease in the right middle frontal gyrus. These regions were positively correlated with increased activity in the episodic memory network [29], activity in specific nodes of this network was associated with better item and context retrieval, have been associated with age-related declines in episodic memory retrieval [30-32], but our patient is young, the mean age is 34.5±9.1 years. So it can infer that HIV infect can accelerate aging. While in other studies it also correlated with the ability to resolve interference efficiently [33, 34]; it reduced will infect selection and inhibition in Chinese semantic judgement.

Meanwhile, we found that, comparing with the control group, bilateral cingulate gyrus volume of HIV patients was the most significantly induced, which is the common changed part in gray matter caused by HIV infection, the result was same with some similar morphologic analysis studies by VBM [27, 35]; the related fMRI...
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studies [36] found that the function of anterior cingulate associated with cognitive was selective attention and conflict management. Then, we found that patient’s gray matter volume decreases lateralization to the left brain, confirms the function of the left brain is closely related to the conscious. Some MRS and MRI results showed HIV-related gray matter atrophy is most common in the basal ganglia (especially the caudate nucleus) [37, 38]. In our study, we did not find the volume of gray matter in the basal ganglia nuclei decreases.

In our study, we also found the right cerebellar crus local volume decreased, as a study of stroke patients showed the cerebellar crus was located in the affected cerebral cortex, which further supports these decreased may represent the impaired cognitive function [39]. So we can infer cognitive impaired from the cerebellar crus local volume decreased.

We found patients’ gray matter volumes around the lateral ventricles and the periaqueductal were increased than the control group. Periaqueductal gray matter (PAG) is constituted with nerve cells gathered around cerebral aqueduct. It is known that PAG functions include the descending modulation of pain, defensive behavior, reproductive behavior and vocal function [40]. Currently there is no literature reported the volume of PAG increase in HIV patients, while there is no literature involved the reason too. We conjectured that it may be the upper neurons damaged leading to the function weakened, and then the lower neurons enhance their function causing gray matter volume increased. The relevant documents can prove periventricular gray matter (PVG) is associated to pain [41] and hypertension [42], the evidence maybe for a non-opioid mechanism [41]. But no evidence support that related to HIV. It is necessary to collect a large number of data to analysis, and proof by clinical pathological anatomy.

As the pretest study showed that female and male patients involved different brain regions of HIV CNS infection. In the further, we will enlarge the sample of female patients to confirm the effect of gender in HIV infection of the CNS.

Conclusion

In this study we observed: Bilateral anterior cingulate cortex, left inferior frontal gyrus triangular part and orbital part, left superior temporal gyrus, right middle frontal gyrus and right cerebellar crus are key brain regions to HIV infection of the central nervous system. In neuroanatomical, the brain structure change can explain some clinical symptoms in patients directing to cognitive impairment. The results of this study provide some evidence of the functional MR imaging for prevention, treatment and assessment of HAND early.

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Disclosure of conflict of interest

None.

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