Reduction of frontal neocortical grey matter associated with affective aggression in patients with temporal lobe epilepsy: an objective voxel by voxel analysis of automatically segmented MRI

Friedrich G Woermann, Ludger Tebartz van Elst, Matthias J Koepp, Samantha L Free, Pamela J Thompson, Michael R Trimble, John S Duncan

Abstract

Background—Interictal episodes of aggression are often reported in patients with epilepsy. Some have characteristics of what has been referred to as episodic dyscontrol or intermittent explosive disorder (IED). Although structural brain abnormalities are thought to play a part in the pathophysiology of aggression, there are few in vivo studies of structural cerebral changes in patients with epilepsy and aggression. Using quantitative MRI, subtle structural brain abnormalities can be investigated in subgroups of patients with both epilepsy and episodes of affective aggression.

Methods—After automated segmentation of cerebral grey matter from T1 weighted MRI, the objective technique of statistical parametric mapping (SPM) was applied to the analysis of 35 control subjects, 24 patients with temporal lobe epilepsy (TLE) with a history of repeated, interictal episodes of aggression, and 24 patients with TLE without episodes of aggression. Both TLE patient groups were compared with each other and with the control subjects on a voxel by voxel basis for increases and decreases of grey matter.

Results—The patients with TLE with aggressive episodes had a decrease of grey matter, most markedly in the left frontal lobe, compared with the control group and with patients with TLE without aggressive episodes.

Conclusion—These findings suggest that a reduction of frontal neocortical grey matter might underly the pathophysiology of aggression in TLE. These voxel by voxel comparisons can guide further in vivo studies into aggression.

Keywords: aggression; epilepsy; voxel based morphometry; magnetic resonance images

Recurrent episodes of interictal affective aggression are a rare but well recognised and reported problem in patients with temporal lobe epilepsy (TLE). F Factors described as being common in patients with epilepsy alone and in those patients with episodic affective aggression alone, were exposure to violence as a child, low socioeconomic status, trauma, infection, and minimal brain damage.

Temporary limbic and frontal lobe structures are involved in the neurobiology of aggressive behaviour. In patients with TLE, we have shown recently, using quantitative MRI, that amygdala or hippocampal sclerosis was not associated with interictal aggressive episodes. This study also showed lower intellect and higher levels of depression and anxiety in patients with TLE to be associated with interictal episodes of aggression and suggested that more widespread, but subtle, cerebral structural abnormalities were a probable aetiological factor.

We have previously used voxel by voxel group comparisons of structural MRI to show differences in cortical grey matter in different epilepsy syndromes.

The aim of the current study was to investigate the structure of the whole cortical grey matter in patients with TLE with or without episodes of affective aggression, using automated grey matter segmentation of T1 weighted MRI images and voxel by voxel comparison between these otherwise clinically homogeneous groups of patients with TLE and normal control subjects.

Methods

CONTROL SUBJECTS AND PATIENTS

Patients with TLE were recruited from a tertiary referral centre (National Society for Epilepsy, National Hospital for Neurology and Neurosurgery). Informed consent was obtained from all patients. A consensus diagnosis of TLE was reached by two neurologists. This diagnosis was based on complex partial seizures with a semiology and EEG findings compatible with a temporal lobe origin in the absence of any EEG or MRI pathology pointing to an extratemporal focus. Patients with an IQ<70 or a history of personality disorder or psychoses were not included.

Twenty five patients with TLE with a history of episodic affective aggression (eight women, six left handers; median age 27 years, range 18–49 years) were identified. These patients fulfilled DSM-IV criteria for intermittent explosive disorder (IED; DSM-IV: 312.34) except that they all had an organic brain disease (TLE). The essential phenomenological feature of IED was the occurrence of discrete episodes of failure to resist aggressive impulses that resulted in serious assaultative acts or...
Table 1: Clinical data in patients with TLE with or without IED

<table>
<thead>
<tr>
<th></th>
<th>TLE with IED</th>
<th>TLE without IED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of onset of epilepsy (median, range; y)</td>
<td>7 (1–28)</td>
<td>8 (1–29)</td>
</tr>
<tr>
<td>Duration of epilepsy (median, range; y)</td>
<td>19 (5–45)</td>
<td>23 (7–46)</td>
</tr>
<tr>
<td>Seizure frequency (mean/month, (SD))</td>
<td>13.4 (15.7)</td>
<td>21 (38.4)</td>
</tr>
<tr>
<td>SDAS 9 (mean, (SD))</td>
<td>14.9 (7.5)</td>
<td>0.4 (0.7)</td>
</tr>
<tr>
<td>SDAS 21 (mean, (SD))</td>
<td>30.9 (12)</td>
<td>3 (4.6)</td>
</tr>
<tr>
<td>FIQ (mean, (SD))</td>
<td>80.6 (8.5)</td>
<td>93 (13.9)</td>
</tr>
<tr>
<td>PIQ (mean, (SD))</td>
<td>81 (8.6)</td>
<td>93.8 (14.1)</td>
</tr>
<tr>
<td>PIQ (mean, (SD))</td>
<td>83.3 (11.8)</td>
<td>94.7 (15.1)</td>
</tr>
<tr>
<td>BDI (mean, (SD))</td>
<td>8.8 (4.8)</td>
<td>4.2 (5.8)</td>
</tr>
<tr>
<td>S-STAII (mean, (SD))</td>
<td>44.1 (14.9)</td>
<td>32.1 (10.9)</td>
</tr>
<tr>
<td>Verbal fluency (mean, (SD))</td>
<td>8.3 (3.7)</td>
<td>12.2 (5.2)</td>
</tr>
</tbody>
</table>

IED=intermittent explosive disorder; SDAS=social dysfunction and aggression scale; FIQ=full IQ; VIQ=verbal IQ, PIQ=performance IQ, all measured with the HAWIE-R; BDI=Beck’s depression inventory; S-STAII= state trait anxiety inventory.

*p<0.01.

destruction of property. The degree of aggressiveness was out of proportion to any provocation or precipitating psychosocial stressors and was not accounted for by another psychiatric disorder. Twenty five patients with TLE without IED were matched for socioeconomic variables, duration of epilepsy, and seizure severity (10 women, two left handers; median age 33 years, range 19–56 years). Thirty five healthy volunteers were scanned (17 women; median age 28 years, range 14–55 years).

The patients with TLE of this current study have been described earlier, using clinical psychiatric and quantitative psychometric assessment as well as visually assessed and rater dependent, volume of interest based quantitative MRI. Clinical and psychometrical data of the patients with epilepsy are summarised in table 1. To assess aggression, carers, and next of kin of both TLE patient groups were asked to fill in the social dysfunction and aggression scale (SDAS-21), including a subscale for outward aggression (SDAS-9). Psychopathology was quantified using well established psychometric questionnaires for depression (Beck’s depression inventory, BDI) and anxiety (state trait anxiety inventory, STAI). Furthermore all patients received extensive neuropsychological testing including an assessment of overall intelligence. A verbal fluency test was included (as the first word of a series beginning with the letter S as possible during a period of 1 minute). Diagnoses on visual assessment of MRI are summarised in table 2.

MRI SCANNING PROTOCOL

Magnetic resonance imaging was performed, using a 1.5 T GE Signa scanner (Milwaukee, USA). An inversion recovery prepared 3D spoiled gradient echo (IRP-SPGR) sequence (TR/TE/TI/NEX 17.4/4.2/450/1, slice angle 20°, matrix size 256x192, 24x18 cm FOV) with 124 contiguous coronal slices and a slice thickness of 1.5 mm was used for volumetric studies. All MRIs were reviewed by experienced neuroradiologists.

Table 2: MRI diagnosis on visual assessment

<table>
<thead>
<tr>
<th>MRI detected abnormalities</th>
<th>Left temporal</th>
<th>Right temporal</th>
<th>Bilateral temporal</th>
<th>No abnormality</th>
</tr>
</thead>
<tbody>
<tr>
<td>TLE with IED</td>
<td>11 (8 HS, 3 others*)</td>
<td>1 HS</td>
<td>3 HS</td>
<td>10</td>
</tr>
<tr>
<td>TLE w/o IED</td>
<td>10 HS</td>
<td>8 HS</td>
<td>1 HS</td>
<td>6</td>
</tr>
</tbody>
</table>

HS=hippocampal sclerosis, *one with amygdala sclerosis, one with low grade glioma of the amygdala, one with dysembryoblastic neuroepithelial tumour of the amygdala; w/o=without.
The resulting significant differences through the 3D image space were displayed collapsed into three orthogonal planes (“glass brain”) (fig 1 A). Regions of significant difference were overlaid on normalised T1 weighted images to facilitate correlation with anatomy (fig 2). The Talairach coordinates\(^1\) are given in mm describing the location of significant voxels; \(x\) defining the lateral displacement of this voxel from the midline (left=negative), \(y\) defining the anteroposterior position relative to the anterior commissure (posterior=negative), and \(z\) defining the vertical position relative to the line connecting the anterior and posterior commissure (down=negative).

**Results**

Combining clinical, electrophysiological, and qualitative MRI information, 14 patients with IED were diagnosed as having left sided TLE, four right sided TLE, and in seven patients with TLE, the seizure onset could not be lateralised using this information (three patients with IED had electrophysiological or imaging signs of bilateral temporal lobe involvement and three had findings equivocal concerning the lateralisation of their TLE). Eleven patients without IED had left sided TLE, nine right sided TLE, and in five patients without IED, the TLE could not be lateralised. None of these TLE lateralisations were overrepresented in either the TLE group with or without IED. One patient from each group could not be scanned for the current study because of a vagus nerve stimulator in one and claustrophobia in the other patient, although previous imaging was available for visual analysis (table 2). Table 2 summarises the MRI diagnoses on visual assessment and shows that amygdala or hippocampal sclerosis in patients with TLE was not associated with interictal aggressive episodes\(^5\); it also shows that right sided hippocampal sclerosis was more frequent in patients with TLE without IED.

Voxel by voxel group comparisons of automatically segmented grey matter were made between 24 patients with TLE each with or without IED and 35 control subjects. In patients with TLE with IED compared as a group with healthy control subjects, reductions of grey matter were found over large areas of the left extratemporal neocortex with maxima in the left frontal neocortex; one maximum difference projection had a Z score of 5.67 at Talairach coordinates \(x=-58, y=36, z=9\) mm (left anterior frontolateral cortex), the other a Z

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**Figure 1** (A) “Glass brain” view of decreased grey matter in 24 patients with TLE with episodes of affective aggression compared with 35 control subjects; displayed after correction for multiple comparisons. The area of maximum difference (Talairach coordinates: \(x=-58, y=36, z=9\) mm) is marked with a white arrow. Another peak area of significantly decreased grey matter (Z score=4.78) was found at Talairach coordinates \(x=-66, y=0, z=28\). (B) Plot of adjusted values of grey matter demonstrates the comparison between the groups in the area of maximum difference (white arrow in A).
score of 4.78 in a more posterior left frontal lobe location (Talairach coordinates x=−66, y=0, z=28 mm, fig 1 A and B). Patients with TLE who did not have IED showed no significant decrease of cortical grey matter compared with control subjects. Patients with TLE with IED also had reduction of left frontal grey matter, compared with patients with TLE without IED (fig 2), although less marked than when compared with control subjects (Z score=3.49 at Talairach coordinates x=−66, y=−2, z=26 mm). The SPM based voxel-wise correlation of SDAS scores and automatically segmented grey matter in all patients with TLE showed a left frontal grey matter area being negatively correlated with these scores which expressed social consequences of interictal aggressive episodes (Z score=3.65 at Talairach coordinates x=−66, y=−2, z=26 mm). Age, scores of depression and anxiety, IQ measures, or scores of verbal fluency did not significantly correlate with specific decreases in grey matter in all patients with TLE.

In patients with TLE with IED compared with control subjects, increases of grey matter were found in the neocortex of the left temporal lobe, whereas similar changes were found bilaterally in patients with TLE without IED (fig 3 A and B).

**Discussion**

This is the first MRI study of a possible structural correlate of interictal aggression or IED in patients with epilepsy, using automated segmentation and voxel by voxel comparisons. We demonstrated a decrease in frontal grey matter in patients with TLE and IED compared with control subjects or with patients with TLE without IED. This finding adds localising information to prior knowledge of widespread, but subtle brain changes in patients with TLE and episodes of aggressive episodes.

**Methodological Consideration**

Although multiple exogenous or endogenous factors may contribute to episodes of aggressive aggression, this paper focuses on the possible underlying structural substrate of this interictal clinical problem in patients with TLE.

The term “episodic dyscontrol” is controversial. There is no such condition included into the ICD-10 and there is still some discussion as to whether or not this dyscontrol syndrome exists as an independent entity. The diagnosis of “intermittent explosive disorder” has been included into the DSM-IV chapter on impulse-control disorders (DSM-IV 312.34). The diagnostic features of IED are basically those of episodic dyscontrol. Using the DSM-IV criteria of IED operationally, we grouped patients...
Figure 3  (A) Areas of increased grey matter in 24 patients with TLE without IED compared with 35 control subjects might represent blurring of the grey-white matter interface ipsilateral to temporal abnormality—that is, widespread gliosis. Both temporal lobes showed abnormalities as the individual temporal abnormalities were evenly distributed between left and right temporal lobe in this patient group (table 2). (B) In patients with TLE with IED, areas of increased extrahippocampal grey matter affected the left temporal lobe. In this group, individual abnormalities were diagnosed more often in the left temporal lobe than in the right (table 2).
with TLE into those with IED and those without. Addressing the measurable social implications of these episodes of aggression, this classification was confirmed subsequently by subscores of validated questionnaires filled in by the next of kin or carers. However, the lack of a patient control group with IED but without TLE reduces the generalisability of our finding of abnormal frontal neocortex beyond our study group.

There have been several studies assessing subtle brain abnormalities in patients with other neuropsychiatric disorders—for example, MRI based quantification using volume of interest approaches in depression and schizophrenia. Voxel by voxel comparisons of neocortical grey matter and sometimes subcortical white matter, which were segmented automatically, have been used in patients with schizophrenia, depression, and Kallmann's syndrome. In patients with epilepsy, this voxel based method has been validated using semi-automated and automated segmentation and added localising information to the finding of widespread structural abnormalities demonstrated with a VOI based approach in patients with epilepsy.

In the current study, T1 weighted images were spatially normalised, automatically segmented, and smoothed before statistical tests were applied to the images on a voxel by voxel basis. The smoothed grey matter images can be thought of as representing the local amount of grey matter. The left frontal reduction of grey matter in patients with TLE and IED might be the imaging correlate of a localised decrease in grey matter volume or neuronal density. Extratemporal grey matter atrophy or neuronal loss in subgroups of patients with TLE, however, are still awaiting pathological confirmation. Hippocampal sclerosis is common in patients with TLE, but not linked to IED. We have shown recently ipsilateral temporal extra-hippocampal, but not extratemporal abnormalities to be specifically associated with hippocampal sclerosis. Here, the same group finding, an increase of grey matter in the temporal neocortex, was demonstrated for both temporal lobes in patients with TLE without IED, possibly reflecting an even distribution of left and right sided hippocampal sclerosis throughout this group. In patients with TLE with IED, there was an increase of grey matter in the left extrahippocampal temporal neocortex, representing a ratio of left to right sided hippocampal sclerosis of 8:1 in this group (table 2). The aetiology of this group finding, a relative increase in temporal neocortical grey matter located at the inside of the automatically segmented grey matter, is unclear. It may represent diminished grey-white matter demarcation in the temporal lobe previously described on T2 weighted images in patients with hippocampal sclerosis and may be the imaging correlate of histological abnormalities including subpial gliosis, subcortical gliosis, microdysgenesis, or differences in cell densities detected in the neocortex of patients with intractable TLE.

PROBLEMS IN THE DIAGNOSIS OF EPILEPSY, INADEQUATE SAMPLING PROCEDURES, SMALL SAMPLE SIZES, LACK OF ADEQUATE CONTROL GROUPS, INADEQUATE MEASURES OF BEHAVIOURAL PHENOMENA, AND CONFONDING VARIABLES SUCH AS INTELLIGENCE HAVE COMPLICATED PREVIOUS STUDIES OF PSYCHIATRIC ABNORMALITIES IN PATIENTS WITH EPILEPSY. Most of these factors have been accounted for in this study. Confounding factors for an MRI study of cerebral grey matter such as age or sex distribution were controlled for by patient selection. There was, however, a difference in the IQ and verbal fluency between patients with TLE with or without IED (table 1). SPM analysis correlating the local amount of grey matter with IQ measures or verbal fluency, however, did not disclose specific changes in grey matter correlated over the range of these neuropsychological measures describing the patient population of this study. Previously, hyperarousal and dyscontrol, both phenomenological aspects of IED, have been associated with low intelligence and with low verbal abilities and were seen as results of subtle diffuse brain damage.

**BIOLOGICAL IMPLICATIONS**

After comparison of automated segmented grey matter on a voxel by voxel basis, patients with TLE with IED had more structural abnormalities outside the affected temporal lobes than patients with TLE without IED. In this study, the reduction of neocortical grey matter was most marked in the left frontal grey matter in patients with TLE and IED, in addition to abnormalities detected on visual MRI assessment which showed a bias towards the left temporal lobe in this patient group (table 2). Whether in patients with TLE, a history of interictal episodes of affective aggression might imply extratemporal abnormalities and leaves these patients less likely to be rendered seizure free by epilepsy surgery needs investigation. The finding of a decrease in left frontal grey matter in patients with TLE with IED is relevant for understanding the neuronal circuitry related to aggression. In the past it has been suggested that abnormalities in cerebral structures which are involved in impulse control and stimulus recognition can lead to sudden outbursts of violent behaviour. Bear suggested that right sided lesions in the so-called ventral system (consisting of inferior temporal cortex, temporolimbic structures, orbitofrontal cortex) could produce a transient release of aggressive responses in animals as long as there was no damage to the right sided so-called dorsal system (inferior parietal, cingulate, dorsofrontal cortex). These ideas have been controversial as many studies support a
more important role of the left hemisphere in aggression.16–18

The reduced frontal grey matter we have shown might be a structural correlate with earlier findings from functional imaging and MR spectroscopy showing a reduced prefrontal glucose metabolism in murderers and significantly lower neuronal markers in the frontal lobes of repetitively violent patients with learning disabilities, although without clear lateralis-
ing effects.37 38 These findings might suggest a localised reduction in frontal grey matter volume or neuronal density to be common to different syndromes involving dyscontrol or aggressive affect.

The amygdala are critical structures within the cortical-subcortical circuits mediating aggressive behaviour, mainly implicated in processing emotional information and in hyperarousal. In animal models, stimulation of the amygdala leads to aggression whereas bilateral lesions render an animal tame. Further frontal stimulation inhibits the amygdala released aggression.39 Thus, at least part of the circuitry of aggression involves the interconnected amygdala and frontal cortices. In our patient groups, the frontal abnormalities might be correlated with dyscontrol, a failure of inhibition of aggressive behaviour. Bouts of aggression have been described as part of the frontal lobe syndromes both in the orbitofrontal (disinhibited) and frontal convexity (apathetic) syndromes.39 Although patients with TLE and IED scored lower on a test of frontal function (verbal fluency; table 1)40 than patients with TLE only, this difference was not correlated with any change in grey matter. Interactions between different brain regions involved in aggressive behaviour and associated additional neuropsychological impairment will have to be clarified in future studies. Voxel by voxel based studies such as ours might guide these future studies by providing results without the bias of operator dependent approaches.

Conclusion

This study suggests a connection between episodes of IED in patients with TLE and reduced frontal grey matter. It is, however, diffic-
ult to infer a causal relation from cross-sectional data, even if our findings resulted from comparisons with a normal and a disease control group. A longitudinal quantitative MRI study in patients with TLE eventually developing episodes of interictal IED would be needed to distinguish aetiological from epilepsy or treatment related abnormalities. The technique used here is widely available, easy to implement, and could be readily applied to other patient groups.

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**NEUROLOGICAL STAMP**

Ilya Metchnikoff (1845–1908)

The Nobel prize in 1908 was awarded jointly to Ilya Metchnikoff and Paul Ehrlich in recognition of their work on immunity. Metchnikoff was one of the most eminent of Pasteur's pupils. He was not medically qualified but taught zoology at Odessa and St Petersburg and from 1873–82 was Professor of Zoology and Comparative Anatomy at Odessa. He was an outspoken Darwinist and in the university’s deteriorating political atmosphere at the time was labelled a “red”. The Tsar’s assassination in 1881 unleashed the reactionaries and antisemites. Metchnikoff tried suicide, on this occasion for the second time by injecting himself with relapsing fever germs. Despite cardiac damage he recovered.

Metchnikoff’s epochal discoveries occurred in late 1882 in Messina, Sicily when during his investigations of digestive processes of larval starfish he noticed that after introduction of a foreign body or bacteria into the body of a transparent starfish, a large number of cells soon surrounded the foreign particle, finally absorbing and dissolving it. He compared the process to the accumulation of white blood cells in human inflammation. This led him to formulate the doctrine of phagocytosis, the destruction of bacteria by white blood cells. He invented the term phagocyte (Gr Phaini, to eat) to connect defence with digestion. The pathologist Virchow encouraged him but warned that biomedical opinion generally would be hostile. Metchnikoff extended his theory by studying the role of phagocytosis in metamorphosis and then infectious disease. The fact that some infected animals succumbed while others survived raised the concept of immunity to infection.

In 1903 Metchnikoff succeeded with Emile Roux in transferring syphilis to apes and also made a preventive mercurous ointment. He also did research on cholera. His later years were largely concerned with the study of the aging factors in man. Methods of inducing longevity were discussed in *The Nature of Man* (1904) and *The Prolongation of Human Life* (1910). He devoted much attention to the question of intestinal sepsis and to the possibility of prolonging life by the ingestion of lactic acid bacilli. To study and control senescence Metchnikoff proposed the establishment of a new scientific discipline he named gerontology. In 1938 the first Congress of Aging was convened in Kiev and in 1962 the USSR Institute of Gerontology was founded. Conferences in the United States in the 1930s led to the formation of the Gerontology Society (1945) and eventually to the National Institute of Aging (1974).

An urn containing Metchnikoff’s ashes was placed in the library of the Pasteur Institute. He has been philatelically honoured by Russia but is shown here with a microscope and the Pasteur Institute on a French stamp issued in 1966 (Stanley Gibbons 1707, Scott B398).

L F HAAS
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