Short report

Tardive dyskinesia: clinical correlation with computed tomography in patients aged less than 60 years

M BRAININ, Th REISNER, J ZEITLHOFER

From the Department of Neurology, Niederösterreichisches Landeskrankenhaus für Psychiatrie und Neurologie Klosterneuburg and from the CT Department, Neurological University Clinic, Vienna, Austria

SUMMARY In a prevalence study of 335 psychiatric in-patients 49 patients (14-6%) with tardive dyskinesia were found. In view of the high prevalence of spontaneous dyskinetic syndromes in elderly patients only patients under 60 years were included (n = 21; mean age: 44-9). Clinical rating was performed with the AIMS-scale. CT measurements of ventricular enlargement and cortical atrophy were obtained. Tardive dyskinesia cases did not differ significantly from healthy controls, though some patients with severe dyskinesia showed signs of brain atrophy. These findings did not provide evidence to support neuropathological reports describing neuronal cell loss and midbrain gliosis in such patients. It is concluded that such structural brain changes in tardive dyskinesia patients are not detectable with present CT technology: they may either be due to concurrent aging processes or, in the case of younger patients, can be confirmed only by more functional methods of testing.

Tardive dyskinesia includes a wide variety of involuntary movements including orolingual dyskinesia, chorea, athetosis, dystonia, tics, and facial grimacing but excluding rhythmic tremor. It occurs in at least 10–20% of the patients exposed to neuroleptic drugs for more than a year. The phenomenology of this disorder has been repeatedly reviewed and expanded over the last few years. A major problem in diagnosis has been the higher prevalence of tardive dyskinesia in elderly patients and the difficulty of determining the prevalence of spontaneous dyskinetic movements in the normal population. Klawans and Barr have recently determined such a baseline, showing that the prevalence of spontaneous lingual-facial-buccal dyskinesias is 0-8% between the age of 50 and 59 years, rising abruptly to 6% and 7-8% within the next two decades.

There have been several neuropathological reports of structural changes in the brain of patients with tardive dyskinesia, especially gliosis and neuronal degeneration in the upper midbrain and in the substantia nigra. However, these cannot be reliably correlated with the clinical disorder. Such changes were attributed to concurrent processes of aging and were therefore not necessarily thought to be a direct effect of prolonged neuroleptic medication.

The present study was designed to answer the questions: do patients with tardive dyskinesia under the age of 60 years show morphological lesions on CT? Are there significant atrophic or gliotic changes compared to controls? Is there a correlation between the clinical severity of abnormal movements and the extent of brain atrophy in such patients?

Material and method

In a prevalence study of tardive dyskinesia in the Niederösterreichischen Landeskrankenhaus Klosterneuburg 335 psychiatric in-patients were examined for symptoms suggestive of tardive dys-
kinesia. Rating was performed by the treating psychiatrists using the AIMS scale. 1 Forty-nine patients with tardive dyskinesia of varying degree were found (14.6%). Out of 132 male patients there were 20 (15%) and out of 203 female patients there were 29 (14.3%).

To exclude possible effects of aging, only patients up to 60 years of age were selected for further study (n = 21; 13 males and eight females). Their ages ranged from 20 to 59 years, the average age being 44.9 years. All patients had a history of neuroleptic therapy of at least two years duration. Consequently, CT examinations were performed with a SOMATOM 2N (Siemens, matrix 256 x 256). At least eight serial slices were made of 8mm thickness. In no case contrast enhancement was applied.

To evaluate atrophic changes three parameters were used. For the lateral ventricles the Ventricle-Index-Ratio (VIR) and the Ventricle-Waist-Ratio (VWR) were used. The VIR is the ratio between the maximum width of the anterior horns and the maximum width of the internal diameter of the skull on the same slice. The VWR is the ratio between the minimal width of the cella media and the maximal internal diameter of the skull on the same slice. VIR and VWR are expressed as percentages. Average values and standard deviations of the VIR and VWR measurements were compared to a normal population under the age of 60 as previously published. 7 In addition, signs of cortical atrophy were evaluated with a simple yes/no judgement by two of the examiners (ThR and MB). Only when found by both examiners cortical atrophic changes were considered valid.

Results

In 21 patients under 60 years of age clinical rating was performed with the AIMS scale and the severity of abnormal movements as well as the severity of oro-facial movements was assessed. CT was performed in 15 of the 21 patients within two weeks of rating. Six patients either refused to have a CT done or were too anxious or psychotic to be included (see table).

Out of 15 patients only five had an abnormal CT scan. In three cases moderate internal hydrocephalus was found (cases 6, 8, 12). In one case moderate atrophic changes were found infratentorially in combination with enlarged cerebro-spinal fluid spaces at the cerebral convexity (case 3). In one case unilateral calcification in the right pallidum was found (case 11). The average VIR (%) was 26.01 ± 4.55 and the average VWR (%) was 18.08 ± 5.16. For average values and standard deviations the r test for independent probes was used, referring to values for normals under the age of 60. 7 No difference was found in VIR and VWR values, respectively (t = 0.011, ns; t = 5.28, ns). Consequently, no relationship could be established between severity of abnormal movements, severity of facial—oral movements and degree of atrophic changes as seen on CT, although four patients with comparatively high clinical scores also had some signs of brain atrophy.

Discussion

CT is an efficient instrument for assessment of diffuse or local atrophic changes in degenerative diseases of the brain. However, atrophic changes of the brain occur with normal aging. Therefore, the diagnosis of “brain atrophy” should not be based solely on CT data, but must include the results of clinical examination. It is also widely acknowledged that the presence of senile dementia of Alzheimer type or multi-infarct type may correlate well with diffuse atrophic changes in the brain as seen on CT, despite a considerable group overlap with controls. 8 There are other data lacking this correlation 9 10 and still others stating a clear-cut difference on CT between patients with dementia and normal controls. 11 Even more controversial is the attempt to correlate the clinical extent of senile dementia with the extent of atrophy seen on CT. With positron emission tomo-
malities included widening in siblings system longer choreoathetosis were CT cortical and means density ganglia basal as a confirmed, although no schizophrenia, ratios and in the copically. '9 the however, graphy, relationship Tardive dyskinesia: Acute and pallidus cerebral cortex.23 olive,22 inferior midbrain and possible to toxic reports of patients, mostly based Neuronal dyskinesia. and occurrence their data changes dive of kinesia of this study. "6 by this attenuation of Wilson's disease clinical studies have been possibly underlie tardive dyskinesia might more clearly be revealed by correlation with the results of functional methods of testing, such as neurophysiological investigations and the evaluation of receptor site dynamics.

References
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