

Extrapyramidal features in advanced Down's syndrome: clinical evaluation and family history

P Vieregge, G Ziemens, M Freudenberg, A Piosinski, A Muysers, B Schulze

Abstract

Extrapyramidal, frontal release, and other neurological signs were studied in 54 demented and non-demented patients with Down's syndrome (DS). Fourteen patients were demented and five showed extrapyramidal signs, mainly of the rigid-hypokinetic spectrum and similar to Parkinsonian features in advanced Alzheimer's disease (AD). None of the non-demented patients had Parkinsonian signs. The mean age of the demented DS patients with extrapyramidal signs was significantly higher than that of the patients without. Frontal release signs were present in demented and non-demented patients. A questionnaire showed no increase in either the proportion of early- or senile-onset dementia or Parkinsonism among first- and second-degree relatives of DS patients. Parkinsonian signs appear to be present at a lower frequency in DS than in advanced AD. A speculative hypothesis about a gene dosage effect of Cu/Zn-superoxide dismutase in preventing toxic radical formation in the substantia nigra of DS patients is presented.

A study of the effects of ageing on individuals with Down's syndrome (DS) may offer a particularly useful high-risk human model for Alzheimer's disease (AD).¹ Several converging lines of evidence support a close association between AD and DS. Neuropathological changes characteristic of AD (neurofibrillary tangles, senile plaques) are found in the brains of almost all DS individuals dying after the age of 35 years.² Moreover, investigators have described age-related abnormalities in visual memory,³ adaptive and social behaviour,⁴ neurological status,⁵⁻⁸ EEG rhythms,⁹ and cerebral glucose metabolism¹⁰ in individual cases or series with DS patients, showing a subtle deterioration with advancing age and similar to the pattern found in AD. Mortality curves for DS individuals between 40-60 years appear remarkably similar to those for the general population between 60-80 years.¹¹

There are, however, discrepancies. Though "Alzheimer-type dementia" is present in some older DS patients, it does not occur with the same frequency as the plaques and tangles, nor is the degree of dementia predicted by the density of plaques and tangles.¹ Furthermore, despite the morphological similarities the genetic relationship between DS and AD remains

unclear. In relatives of index cases with AD a raised frequency for DS, AD, and haematological malignancies has been found,¹²⁻¹⁴ and an increased number of pre-senile dementia in relatives of DS probands was reported.¹⁵

There is growing evidence that patients with AD of early onset and familial occurrence may have a predisposing locus on chromosome 21, while in families with senile onset such linkage could not be substantiated.^{16,17}

Patients with AD of an advanced stage exhibit extrapyramidal (EP) and particularly Parkinsonian signs to various degrees.¹⁸⁻²⁷ A recent case control study provided evidence for a familial aggregation of AD with Parkinson's disease (PD).²⁸ At necropsy, cell loss, neuromelanin deposition, Lewy body formation, and gliosis in the substantia nigra (SN) has been demonstrated in a proportion of AD brains.^{26,29,30} Extrapyramidal signs in DS have until recently been referred to in one larger study³¹ and in single cases.^{10,32-34}

To clarify the natural history of DS patients in middle and old age, we looked for EP system involvement and other neurological signs in DS patients and for evidence of familial clustering of AD and PD in families with an index case of DS.

Patients and methods

Fifty four patients (34 males, 20 females) with DS from various institutions were included in the study. The diagnosis of DS was established on the basis of the classic phenotype. Karyotype analysis was carried out wherever possible: 27 patients had "free" trisomy 21, two had a mosaic pattern, and one a translocation trisomy.

The history of each patient was taken. Special attention was given to diseases known to occur more frequently in DS, such as cardiac anomalies, myeloproliferative disorders, thyroid disease, previous neurological diseases and medication. The degree of characteristic phenotype of DS was assessed clinically. A neurological examination as detailed as possible in the individual patients was performed personally by one of us (PV). Fixed criteria were used according to the Unified Parkinson's Disease Rating Scale (UPDRS)³⁵ to evaluate the degree of hypokinesia, rigidity, resting tremor, postural abnormalities, walking pattern, and pulsion phenomena. Special attention was given to dyskinesias, myoclonic jerks, pyramidal signs

Medizinische
Universität zu Lübeck,
Lübeck Klinik für
Neurologie
P Vieregge

Institut für
Humangenetik
A Muysers

Landeskrankenhaus,
Heiligenhafen
G Ziemens

Landeskrankenhaus,
Neustadt/Holst
M Freudenberg
B Schulze

Psychiatrisches
Krankenhaus,
Rickling, FRG
A Piosinski

Correspondence to:
Dr Vieregge, Klinik für
Neurologie, Medizinische
Universität zu Lübeck,
Ratzeburger Allee 160,
D-2400 Lübeck 1, FRG

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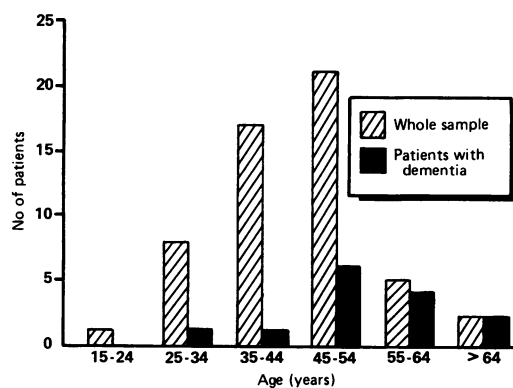


Figure Age distribution of 54 patients with Down's syndrome evaluated for extrapyramidal signs and dementia.

and signs of "frontal disinhibition" such as palmonental reflex (PMR), grasping reflex (GR), sucking and/or snouting reflex (SSR).

Dementia was clinically assessed by interview and/or by assessment of those caring for the patients using items about deterioration of personality, loss of initiative and conversation, self-help skills and cooperation, memory deficits, concentration, disorientation. Psychometric tests were not applied. Thirty five patients had EEG examination, nine patients cranial CT. A postal questionnaire on the occurrence of dementia and Parkinsonism in first- and second-degree relatives was obtained from the families of 38 patients.

Three patients (aged 17, 43, and 50 years) died within weeks or months of the clinical examination. At necropsy, both of the two older patients displayed typical brain morphology with atrophy of the superior temporal gyrus macroscopically and diffusely spread senile plaques and neurofibrillary tangles in the cortex microscopically.

Results

The age distribution of the whole patient sample is given in the figure. Table 1 shows the mean ages of the patient sample for dementia, extrapyramidal, and frontal signs. The mean age between female and male patients did not differ significantly. Fourteen DS patients (age range 27-67 years) were assessed to be demen-

Table 1 Number and mean (SEM) age of patients with Down's syndrome with dementia, extrapyramidal involvement, and signs of frontal disinhibition

Patient group	Number (%)	Mean age (years)
Whole sample	54	44.4 (9.9)
Male patients	34 (63)	44.4 (10.0)
Female patients	20 (37)	44.9 (10.0) NS
Non-demented	40 (74)	42.2 (8.6)
Demented	14 (26)	50.9 (10.9) ***
Extrapyramidal signs present	5/14 (36)	59.8 (6.7) **
absent	9/14 (64)	45.8 (9.7)
PMR present	30 (56)	48.7 (8.1) ***
absent	24 (44)	39.0 (9.5)
GR present	5 (9)	54.2 (11.1) *
absent	49 (91)	43.4 (9.4)
SSR present	4 (7)	47.3 (3.8)
absent	50 (93)	44.2 (10.2) NS

*, **, *** denotes $p < 0.05$, < 0.01 , < 0.001 respectively (Student's *t*-test); NS=not significant; PMR=palmonental reflex; GR=grasping reflex; SSR=sucking and snouting reflex.

ted. Their mean age was significantly higher than that of the 40 non-demented DS patients.

The mean age of five demented DS patients with extrapyramidal signs was significantly higher than that of the nine demented patients without extrapyramidal involvement. Signs of frontal disinhibition were present in demented as well as in non-demented DS patients. The mean age of patients with palmonental reflex and of those with grasping reflex was significantly higher than that of the patients without these signs. In the extrapyramidal involvement rigid-hypokinetic signs prevailed (table 2). Two further patients had oral dyskinesias, but were not included in the study as they were on current neuroleptic medication. Resting tremor and myoclonic jerks were not observed. No patient received levodopa treatment.

Other neurological signs unrelated to dementia were strabismus ($n = 17$), nystagmus ($n = 3$), pyramidal signs ($n = 3$), dysarthria ($n = 1$). Precocious ageing signs were: early grey hair ($n = 5$), cataract ($n = 3$), hypacusis ($n = 2$). Grand-mal epilepsy was present in seven patients (two demented); two patients had occasional seizures. Seizures were observed a few weeks before death in one patient. Two patients were on anticonvulsant medication. EEG was normal in 11 patients, mildly abnormal in 13, and moderately abnormal in 11. None of the patients with seizures had epileptic discharges on the interictal EEG recording. There was no relation between severity of EEG changes and signs of frontal disinhibition or dementia.

Atrophic changes on CT were seen in two demented patients. Three non-demented patients had basal ganglia calcification, but no signs of extrapyramidal involvement.

No first-degree relatives with early-onset dementia (before 65 years) could be detected from the questionnaire. One mother had Parkinsonism before the age of 65. Among 149 first- and second-degree relatives older than 65 years two had senile dementia (mother, paternal grandfather), and two had Parkinsonism (father, paternal grandfather). Other single neurological disorders in first-degree relatives were essential tremor, glioma, meningioma, epilepsy, and DS (a brother).

Discussion

Conclusions from our study should be drawn cautiously because of the methodological limitations in the clinical investigation of men-

Table 2 Extrapyramidal and "frontal" signs in 14 demented patients with Down's syndrome

Sign	Number of patients (%)
Rigidity	3 (21)
Shuffling gait	4 (29)
Hypokinesia	3 (21)
Hypomimia	3 (21)
Orofacial dyskinesia	2 (14)
Palmonental reflex	9 (64)
Grasping reflex	3 (21)
Sucking/snouting reflex	2 (14)

tally handicapped persons. In DS, nonspecific neurological signs can be observed at any age group.^{5,6} The frequency of frontal release and pyramidal signs, increased muscle tone, and hyperreflexia rises with advancing age.^{5-8,36} In our investigation only the palmomental reflex was more frequently seen in the older patients. In contrast to the literature, other frontal release signs were less frequent and were also observed in non-demented DS patients.^{6-8,31}

Since our investigation was cross-sectional and not prospective, it may have missed some neurological signs eventually occurring later in the course of advanced DS. This may explain the lower frequency of seizures and lack of epileptic discharges among the nonspecific EEG changes in our patient sample compared with other reports.^{7,8} Recent longitudinal studies have observed pre-terminal myoclonic jerks and seizures quite regularly in the course of advanced DS.^{31,37} To a lesser degree such myoclonic and seizure activity is also found in advanced AD.²⁴

No control group was used in this study because both the rate of development and the extent of extrapyramidal signs in DS was unknown and DS is a neurologically unique model of precocious ageing. We expected that the extrapyramidal system involvement might be encountered in DS patients with dementia, the latter most obvious on clinical presentation. Psychometric testing was therefore not applied. In this way the study confirmed that at a given stage of middle or old age only a certain percentage of DS patients are demented (26% in our sample).

Extrapyramidal signs in older DS patients (36% of demented; 9% of all) were more frequently seen than in other reports, where such signs were not mentioned or even absent.^{5,6,8,36} Lott *et al* refer to "gait deterioration" in 73% of their middle and old-aged patients late in the disease course.⁷ In a most recent investigation 20% of demented DS patients of middle and old age showed Parkinsonian signs.³¹ These different results may be due partly to the fact that the age group above 50 years is underrepresented in some studies.^{5,6} Our study was focused on EP signs and fixed criteria were used for EP evaluation.

Extrapyramidal signs in DS have up to now been described in single cases, for example, a 47 year old woman had choreiform movements and at necropsy had the morphological features of Pick's disease together with atypical AD;³³ a demented 63 year old DS patient had gait difficulties, rigidity, and nonspecified tremor.¹⁰ Postural tremor in advanced DS has been described in a patient by Jervis;³² a 23 year old male DS patient had developed resting tremor two years previously and showed masked face, cogwheel rigidity, generalised hypertonia, and the unsteady, festinant gait with reduced arm swing.³⁴

Resting or postural tremor was not observed in our sample, but had been present in four out of 10 DS patients with EP signs in another larger sample on dementia in DS.³¹ Two patients of that prospective investigation had Parkinsonism before the onset of dementia,

whereas all of our DS patients with EP signs were demented. These patients were of a significantly higher age compared with those demented DS patients without EP involvement. It appears therefore from our study, that EP signs are features more likely to occur late in the clinical course of demented DS patients, but that such signs do not seem to be essential in the disease spectrum.

We found 36% of demented patients with EP signs in DS are in the range also found in AD patients.²³⁻²⁶ Others have described different percentages for their AD patients (9%,²¹ 62%,¹⁸ 65%,²⁷ 92%²²).

The main clinical feature in our study was of the hypokinetic rigid type. This bears similarities to Parkinsonism encountered during the course of AD.¹⁸⁻²⁷ In our DS series none of the patients had tremor at rest, which in AD has been observed in between 4-48%.^{18,20,22,25,27} Dyskinesias in AD unrelated to neuroleptic medication have been seen in 17%,²² 23%,²⁰ and 38%.²⁵ In an unselected population of non-demented older people the rate was 38%.³⁸ In order to clarify, if dyskinesias in DS are more frequently present than 14% (our series), a prospective lifetime study of the individual patients seems necessary.

The occurrence of EP signs in AD can be sufficiently related to pathological changes in the substantia nigra (SN).^{26,29,30} Also in DS, several neuropathological and neurochemical theories can support the clinical findings.

First, it is obvious from ours and from previous studies, that calcification of the basal ganglia, which is frequently encountered in DS, does not play a pathogenetic role in the development of EP features.³⁹

The SN is less severely involved pathologically than other subcortical nuclei in young and middle aged DS patients, in contrast with AD.^{40,41} On the other hand, the brains of patients with advanced DS show more numerous neurofibrillar tangles than comparable AD brains and mild to moderate SN pathology suggesting Parkinsonism (Lewy bodies, neuromelanin deposits).⁴² Others, however, did not observe Lewy bodies in older DS patients.⁴³ Neurochemically the content of dopamine in the caudate nucleus and of homovanillic acid in the CSF is reduced in more advanced stages, while it is normal in young and middle aged DS patients.^{40,44,45}

Together, these findings support the finding that the pathological changes in the SN and the clinical presentation of Parkinsonism seem to be phenomena occurring late in the course of DS.

Why does the SN in DS seem to be afflicted so late in the disease compared with other regions of the brain and to a less severe degree than in AD? One should remember that the triplication of the chromosomal segment 21q22 is responsible for an overexpression of the gene for the copper/zinc-superoxide dismutase (Cu/Zn-SOD). Such elevated gene dose may disturb the steady-state of active oxygen species within the cell, resulting in oxidative damage to biologically important molecules. It has been suggested that such a mechanism may in part

be responsible for the mental retardation and for the AD pathology (that is, paired helical filament formation) associated with the DS phenotype.⁴⁶

Cu/Zn-SOD on the other hand catalyses the conversion of superoxide radicals to hydrogen peroxide. In Parkinson's disease hydrogen peroxide and related oxygen radicals may be involved in the degeneration of dopamine neurons.⁴⁷ Cytosolic Cu/Zn-SOD-like activity and mitochondrial Mn-SOD-like activity are increased in the SN of Parkinsonian brains at necropsy.^{48,49} In DS patients perhaps the excess of Cu/Zn-SOD serves as a protective mechanism in the SN by scavenging radicals occurring in nigral metabolism. This may result clinically in a Parkinsonian syndrome at a later stage and, possibly, to a lesser degree than expected from comparison with AD. It has been also shown that Cu/Zn-SOD protects against toxic effects of pyridinium derivatives (for example, paraquat), that are currently considered to play a pathogenetic role in Parkinsonism.⁵⁰

It may also be possible that overexpression of Cu/Zn-SOD in familial, early-onset AD could result in Parkinsonian features at a lesser degree and, probably a later stage, when compared with those AD cases without linkage to chromosome 21.

As mentioned earlier, there is a clinical and pathological overlap mainly in the late stages in AD and PD. A recent family study reported an increased risk for AD and for PD among the relatives of probands with AD.²⁸ Previous pedigree analyses of index patients with AD suggesting an increased risk in their relatives to suffer from AD, DS, and/or haematological malignancies have been criticised mainly on grounds of methodology and sample size.^{12,14} As a raised frequency of presenile dementia in relatives of DS patients has been reported,¹⁵ we undertook a similar, questionnaire-based inquiry on dementia and additionally on PD signs in the families of our DS patients. There was no increased rate of dementia with onset before the age of 65 years. Two senile-onset demented relatives among 149 persons at risk does not greatly exceed that expected from the age-specific prevalence rate for dementia over 65 years of age in the general population (0.65%),⁵¹ (2×100): 149 = 1.34%. The same applies for two relatives with senile-onset (after age 65 years) PD, given the similar age-specific prevalence rates for dementia in the seventh and eighth decade of life.

Thus our data do not provide better evidence of the relative risk of AD and PD in families with DS probands. This is mostly because the sample is small, the method was questionnaire-based, and diagnostic proof of the cases at risk was not possible.

This investigation has shown that extrapyramidal signs may be a feature of advanced DS. They were encountered only in demented DS patients. This finding parallels Alzheimer's disease, where such signs may also occur in an advanced stage, but to a greater degree. To evaluate the extrapyramidal involvement in DS may help to clarify the debated association between AD and PD, and further clinico-

pathological and neuro-chemical studies in DS patients may be useful. Such studies may gain additional interest considering the role of free-radical formation and its prevention in the pathophysiology of idiopathic Parkinson's disease.

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