Smoking and dementia of Alzheimer type

Sir: Several surveys have reported a negative association between smoking habits and Parkinson’s disease suggesting that smoking may decrease an individual’s risk of developing the condition. An observation made by Appel that only six of 30 Alzheimer patients in his study had smoked at any time in their lives, led him to conclude that a similar effect may operate in dementia of Alzheimer type (DAT). This is of interest in view of the recently demonstrated loss of nicotinic receptors in the cerebral cortex of DAT patients.

Conversely, Shalat has presented data from a case control study, showing that Alzheimer patients were more than twice as likely to be smokers or ex-smokers than were controls. Furthermore, increased risk of the disease was positively correlated with level of cigarette consumption.

Our data refer to patients attending the Maudsley Memory Clinic and the psychogeriatric unit of the Royal Bethlem and Maudsley Hospitals. The medical notes of 81 persons over the age of 65 with a diagnosis of “probable” dementia of Alzheimer type were reviewed to obtain information about smoking habits. Where such information was not recorded the next of kin were contacted by telephone. One hundred and twelve persons from a local luncheon club and care home were selected as age-matched controls and interviewed.

Seventy one per cent of male DAT patients and 32% of female DAT patients either smoked or gave a history of having smoked, as compared with 77% of male controls and 44% of female controls. Even allowing for sex differences (with the use of an appropriate prop-con-linear model) no significant difference in the lifetime prevalence of smoking between the two groups was found (chi² = 1.23, p = 0.05, Odd’s ratio = 1.58).

Our results do not support either study referred to above but are in accord with the findings of an epidemiological study of patients developing the disease before the age of 70.

Clearly further studies are required to elucidate this question but can only be conducted if past and present smoking behaviour is perceived as an important variable and therefore accurately recorded in medical notes.

GEMMA MM JONES
MARGARET REITH
MP PHILPOT
BARBARA J SAHAKIAN
Section of Old Age Psychiatry, Institute of Psychiatry,
De Crespigny Park,
London SE5 8AF, UK

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and slight subjective hearing impairment on the left. Tracheostomy was performed on the third hospital day and IPPV was continued for two weeks. During subsequent weaning from the respirator his breathing was maintained while awake, but during sleep there were frequent episodes of central apnoea with arterial oxygen saturation frequently dropping to 85%.

Neurological improvement began two weeks after admission. At three months the major findings included continuing apnoeic episodes while asleep requiring nocturnal ventilation, complete left facial paralysis, markedly impaired swallowing, mild incoordination of the left arm and leg and mild unsteadiness of gait.

At the end of the fifth month there was development over five days of marked unsteadiness of gait, inco-ordination of the arms and increasing difficulty in swallowing. A CT brain scan and CSF examination were normal. Recovery to his previous state occurred over the next 6 weeks.

At the end of 8 months assisted respiration was no longer required during sleep and the tracheostomy had been closed. There were no significant changes during the next three years.

Somatosensory evoked potentials (SEPs) and brain stem auditory evoked potentials (BAEPs) were recorded during the acute illness and the recovery period using standard methods and normal values for this laboratory. On the second hospital day SEPs were normal but BAEPs following stimulation of the right ear showed slight prolongation of the Wave I-V interval. Nine days later SEPs remained normal but the BAEPs showed a further increase of the Wave I-V interval on the right and loss of all waves following Wave I on the left, despite stimulation at least 55 decibels above left ear hearing threshold. On hospital day 29, while the patient was recovering, the central somatosensory conduction time to the left hemisphere was relatively prolonged compared with that to the right. Subsequently the central conduction time to the right hemisphere became absolutely prolonged while that to the left remained unchanged. Although the BAEPs from the right ear returned to normal, those from the left beyond Wave I remained absent until recordings made at the time of deterioration in the 5th month, when they had returned but showed slight prolongation of the I-V interval. Thirty-two months after admission SEPs and BAEPs were normal.

This patient's presentation with a systemic prodromal illness followed by rapidly progressive multifocal signs in the brain stem was similar to those previously reported in the literature. As in most of these cases, the organism was isolated from blood cultures but not from the CSF. The loss of automatic respiratory movements (Ondine's curse) has not been described in previous patients with listeria rhombencephalitis, although "respiratory failure" has occurred in two. This unusual but important complication might be expected in patchy disease of the lower brain stem from involvement of central respiratory control centres.

The evoked potential studies in this patient showed several unexpected features. The disappearance of all BAEPs beyond Wave I is consistent with a pontomedullary lesion; although this change with preservation of hearing may be seen in multiple sclerosis (unpublished observations) we have not observed it in any other neurological diseases. The later deterioration in the central somatosensory conduction time to the right hemisphere, occurring while the patient was recovering, suggested the development of a new subclinical brain stem lesion. Following this, another lesion in the brain stem-cerebellar connections occurred at five months causing inco-ordination and ataxia.

Little is known of the histopathology of listeria rhombencephalitis. Formation of multiple small abscesses with gram positive bacilli has been reported at necropsy and the evidence suggests that direct bacterial invasion occurs in the acute stage. There is an analogy in listeria infection of rumi

Asymmetry of pathology in Alzheimer's disease

Sir: For much recent work on neural transmitter abnormalities in Alzheimer's disease it has been the practice in many centres to fix one cerebral hemisphere for histological study and to freeze the other for biochemical investigation. The assumption underlying this is that the disease process affects the brain symmetrically, allowing correlation to be made between histological changes on one side and chemical changes on the other. This assumption has been called into question in a recent study by Arendt et al. These authors reported cell counts in the nucleus basalis and plaque counts in the cortex on both sides of the brain in cases of Alzheimer's disease. They found, in some cases, "marked differences in regional plaque counts between the two hemispheres". These differences in the mean counts for the hemispheres did not reach statistical significance, however. Nevertheless, we felt that the doubt cast on the current practice of dividing the brain sagittally called for a review of other data in the literature and a small additional study designed to address...