Motor neuron disease and polio in Scotland

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Abstract
An analysis of mortality and morbidity rates for motor neuron disease (MND) in Scotland has confirmed earlier observations that the disease is more common in men and older age groups. The geographical distribution is non-uniform and related to discharge rates for all neurological diseases. Discharge and mortality rates are increasing but there has been no decline in populations who would have been vaccinated against polio.

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The general term motor neuron disease (MND) refers to those conditions characterised by progressive muscle wasting and weakness caused by a variable combination of upper and lower motor neuron degeneration. Five to 10% of cases are familial and in some pedigrees there is evidence of a gene conferring disease susceptibility located on chromosome 21. Most cases are sporadic, however, and the low concordance rate demonstrated in twin studies provides strong evidence for an exogenous aetiology in this group. In most patients there is no identifiable cause, and although toxic, metabolic, traumatic, infectious and immunological mechanisms have come under close scrutiny, the aetiology is unknown.

There has been longstanding interest in viral hypotheses and the role of poliovirus has received most attention. Charcot was the first to propose a relationship between poliomyelitis and the later development of progressive muscular atrophy in the discussion of a case reported by Raymond and the association has since been described in several case series. Evidence for an association is conflicting, however, and most patients have no history of this condition. Nevertheless, polio infection is usually asymptomatic and argued that, even in patients with no history of poliomyelitis, subclinical infection might be severe enough to deplete motor neurons and that MND might then develop as a consequence of further motor neuron loss due to ageing or a second insult. If this hypothesis is correct, the epidemiology of polio and MND should be related and, in this study, a comparison was made of Scottish mortality and morbidity returns for both conditions.

Method
Mortality data
Since 1968, information about patients discharged from Scottish hospitals has been entered on computers by the Common Services Agency for the purposes of the Scottish Hospitals In Patient Survey. Entries for patients discharged with a diagnosis of motor neuron disease (International Classification of Diseases, eighth revision (ICD8) code 348 from 1965 to 1978 and ICD9 code 335 from 1979 onwards) were retrieved for the period 1968-87 and the data were indexed by name and date of birth to link the records and identify individuals who were discharged with this syndrome for the first time. Discharge rates for all neurological diseases (ICD 320-389) were obtained from Scottish hospital inpatient statistics and these included first and subsequent discharges. Data on the uptake of polio vaccines and notification rates for poliomyelitis were obtained from Scottish Office publications for the period 1931-87.

Mortality data
Mortality statistics for MND, poliomyelitis and other neurological diseases were obtained from publications of the Registrar General for Scotland. Mortality returns for progressive muscular atrophy were first made in 1931 (ICD4 code 81.1 from 1931 to 37). In 1938 the classification was widened to include motor neuron disease (ICD5 code 82 from 1938 to 1947). Since then the categories have been slightly modified (ICD6 and ICD7 code 356 from 1948 to 1964, ICD8 code 348 from 1965 to 1978) and in the most recent revision, MND was classified with other anterior horn cell diseases (ICD9 code 335 since 1979). The classification of acute poliomyelitis has remained reasonably constant since 1929 (ICD4 code 16, ICD5 code 36, ICD6 and 7 codes 80, ICD8 code 40, ICD9 code 45). The same source was also used to obtain mortality returns for all neurological diseases (ICD9 320-389) between 1968 and 1987.

Demographic data
Population estimates for Scotland were obtained from the annual reports of the Registrar General for the years 1931-87 and additional demographic data were obtained from the county census reports for 1931 and small area statistics from the 1981 census.

Age–sex distribution
Age–sex–specific first discharge and mortality rates for MND were calculated for the period 1968-87, using the 1981 population as the denominator. Poisson tables were used to estimate 95% confidence intervals.
Temporal variations in mortality and morbidity

Temporal trends were analysed by determining the mortality rate for poliomyelitis, progressive muscular atrophy and motor neuron disease between 1931 and 1987. The population estimates for the first year of each quinquennium for this period were used as the denominator. Age-specific first discharge and mortality rates were also calculated for the 20-year period from 1968 to 1987 using the annual estimate of the usually resident population as a denominator. The age-sex-specific rates for 1968 to 1987 were then used to calculate annual standardised discharge and mortality ratios for this 20-year period using the method of indirect standardisation (Scotland 1968–87 = 100). The statistical significance of the observed temporal trends was examined using Poisson regression techniques to calculate the average proportionate change in mortality and morbidity per year, together with 95% confidence intervals.

Geographical variations in mortality and morbidity

Crude mortality rates for poliomyelitis and MND (1931–49) were calculated for the 33 counties of Scotland using 1931 estimates of the population as a denominator. These were compared with crude mortality and discharge rates for MND between 1968 and 1974, adjusted to estimates of the population resident in 1968. In 1974 the administrative boundaries of Scotland were changed and the 33 counties were reorganised into nine regions, three island areas, 53 districts and 1211 postcode sectors. Crude mortality rates for regions, island areas and districts were calculated for the period 1975–87 by using the 1981 estimates of population as a denominator. Postcodes were used to identify the area of residence of patients discharged between 1975 and 1987 and, using these data, it was possible to calculate crude discharge rates to postcode sector level for the same period. Age-sex standardised discharge and mortality ratios for regions and districts were then calculated by the indirect method (Scotland 1975–87 = 100). Confidence intervals were calculated using tables of the Poisson distribution and the results were mapped to postcode sector level using Linemap, the computerised mapping programme.

Results

Age-sex distribution

Between 1968 and 1987, 6009 patients were discharged from public Scottish hospitals with a diagnosis of MND and the data were linked to identify 3802 individuals who were classified as having this syndrome for the first time. The mean discharge rate was 3.69/10^5 per year and the disease was more common in men (4.33/10^5 per year) than women (3.09/10^5 per year). The mean age of discharged patients was 61.9 years (SD 16.6). There were 2153 (56.6%) men and they were slightly younger (60.7 years, SD 16.3) than the women (63.4 years, SD 16.7).

Progressive muscular atrophy or motor neuron disease was recorded as the underlying cause of death in 4272 registrations between 1931 and 1987, and 1673 of these returns were made between 1968 and 1987. Age-sex-specific discharge and mortality rates are shown in figure 1. The mean mortality rate was 1.62/10^5 per year and again the mortality rate was higher in men (1.79/10^5 per year) than women (1.47/10^5 per year). The disease was more common in men in all age groups and incidence increased with age. Peak morbidity was observed in 80–84 year olds with mortality becoming maximal in the 65–69 year age group.

Temporal variations

The Salk vaccine was introduced to Scotland in 1956 and surveys in 1961 indicate that 48% of the population born between 1933 and 1942 and 85% of those born between 1943 and 1961 were vaccinated. The live (Sabin) vaccine was introduced in 1962 and infant vaccination rates of 85% have been recorded since. These measures have been accompanied by a marked decline in polio notifications and mortality (table 1) from a peak of 114 and 9.6/10^5 per year respectively between 1946 and 1950 to virtually zero in recent years.

Between 1968 and 1987, crude first dis-
Over the same 20 year period, crude mortality rates for MND also increased significantly from 1·25 to 2·1/10^5 per year (p < 0·05) and age-specific mortality rates rose significantly (p < 0·05) in the 0–19 and 40+ year age groups (table 2). Age–sex standardised mortality ratios increased significantly from 78 to 124 (p < 0·05) and, whilst the number of patients dying from all neurological diseases increased, the proportion with MND still rose significantly from 9·5 to 13% (p < 0·05).

### Geographical variations
Mortality from progressive muscular atrophy (1931–49) was significantly higher in rural areas (r = 0·51; 95% confidence interval 0·20 to 0·73) and there were negative associations between mortality rates for polio for this early period and more recent first discharge and mortality rates for MND for the period 1968–74 (r = −0·48; 95% confidence interval −0·71 to −0·21; and r = −0·21; 95% confidence interval −0·52 to 0·15 respectively).

The postcodes of area of residence were available for 2403 (92%) of the MND patients hospitalised for the first time between 1975 and 1987 (table 3, figure 3). Significant variations were observed with standard discharge ratios ranging from 34 in the Fife region to 119 in the Borders region. Standardised mortality ratios were also non-uniform with values ranging between 77 in the Fife region to 156 in the island areas. Standardised first discharge ratios for MND were inversely related to population density (r = −0·31; 95% confidence interval −0·53 to −0·05) and strongly associated with discharge rates for all neurological conditions (r = 0·78; 95% confidence interval 0·38 to 0·94).

### Discussion
Variations in the incidence of MND should provide clues to underlying aetiology, but it is difficult to measure this statistic because the condition is rare and the onset is insidious. In the United Kingdom morbidity and mortality returns are collected routinely by government agencies and these should provide some information about the frequency of this disease. Holloway has observed that first discharge data tend to overestimate incidence by as much as 100% because of false-positive coding and diagnosis, and mortality returns tend to under represent incidence because of non-certification in 20–30% of cases. Despite these problems morbidity and mortality provide upper and lower limits of incidence so it was possible to use these statistics to test the hypothesis that the distribution of poliomyelitis and MND are related.

### Age-sex distribution
MND morbidity and mortality increased with age and were more common in adult males. The slight decline in the older age groups is probably related to underascertainment. Between 1975 and 1987, 295 (11%) of the first discharge registrations and 88 (7·3%) of mortality returns related to individuals younger...
than 45 years. Whilst it was not possible to determine the vaccination status of these patients, Scottish office statistics indicate that 85% of the population born since 1943 have been vaccinated against polio.

### Temporal trends

In an earlier study from Scotland, Holloway suggested morbidity and mortality from MND were increasing. Our study confirms her earlier findings and similar trends have been observed in England and Wales, the United States and Norway. Swash has argued that these temporal trends may be explained by the rising number of medical practitioners able to make the diagnosis. But, whilst it is true that the number of neurologists has increased, especially in the United States, there has been no comparable increase in the frequency of Parkinson’s disease in that country and the improvements in neurological services in the United Kingdom and Scandinavia have been more modest. Moreover, as the proportion of neurological patients dying from MND has increased significantly it would seem likely that the observed trends reflect a real increase in underlying disease frequency with some additional inflation due to superior case ascertainment and changes in disease classification.

These observations would obviously lend support to the hypothesis that MND is caused by an exogenous factor and, if the polio hypothesis is correct, it is possible to predict a rise in disease frequency, reflecting changes in the incidence of poliomyelitis that occurred in the earlier part of the century, followed by a decline after 2010 as the effects of vaccination become apparent. Epidemiological studies of serological status indicate, however, that antibodies may be acquired in the second decade of life and, if such at-risk populations were successfully vaccinated in the 1960s, one would expect to see an effect on morbidity and mortality as these patients enter the third and fourth decades in the 1980s. But, although polio mortality and notification rates have declined to vanishingly low levels, these changes have not been accompanied by any significant reduction in MND mortality or morbidity in populations who have been vaccinated.

### Geographical variations

In this study there was no evidence of a geographical association between past mortality from poliomyelitis and present morbidity or mortality from MND. It was not possible to replicate Martyn’s study exactly because polio notification rates were not available on a regional basis. In an earlier study from Scotland, Holloway observed that the disease was more common in rural areas but, while this was also true of the present study, standardised first discharge ratios were more strongly correlated with discharge rates for all neurological diseases, indicating that the observed trends might be related to variations in ascertainment.

To conclude, we have found no evidence that MND is caused by early asymptomatic polio infection or that polio vaccination offers protection against this condition. However,
these results must be interpreted with caution because the population is small and both geographical and temporal trends can be biased by variations in case ascertainment and data collection. It will therefore be of critical importance to monitor the incidence of MND in vaccinated populations and perform case-control studies to examine social conditions and vaccination status in early life.

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