Evidence of underdiagnosis of myasthenia gravis in older people

A Vincent, L Clover, C Buckley, J Grimley Evans, P M Rothwell, and the UK Myasthenia Gravis Survey

Background: Myasthenia gravis is a potentially serious but treatable muscle disease caused by autoantibodies directed at the acetylcholine receptor (AChR) on the postsynaptic membrane of the neuromuscular junction. It is generally thought to affect around 10–15 per 100,000 of the population and to be most common in young women. Such patients present during early adult life, often have enlarged thymuses with numerous germinal centres, tend to improve after thymectomy, and are likely to have the HLA B8, DR3 antigen. Less commonly (< 10% of the total), the disease occurs in patients with thymic tumours.

Myasthenia gravis can also present in middle age or later. In this situation, the thymus is not usually abnormal, there is only a weak association with HLA B7 DR2 antigen, and the sex ratio is nearer unity. It is now recognised that the incidence and prevalence of myasthenia gravis in middle aged and older patients are greater than was previously thought. However, the clinical features tend to be different from those in younger cases, with more bulbar involvement in older people, and the differential diagnosis is wider. Consequently, myasthenia gravis in later life is sometimes mistaken for other conditions, such as stroke, motor neurone disease, or parkinsonism, and may be underdiagnosed.

In two previous studies, an apparent fall in the incidence of clinically recognised myasthenia gravis in people over 80 years of age in our national AChR antibody incidence study, and the high prevalence of previously unrecognised positive AChR antibodies in those ≥ 75 years old, suggest that myasthenia gravis may be substantially underdiagnosed in older people.

METHODS

Assay data

We obtained data on 3836 individuals who tested positive for AChR antibody at the nine British centres that are registered with EQAS during three consecutive years (1997 to 1999). All use a standard immunoprecipitation test with [125I-α-bungarotoxin labelled human AChR. These laboratories perform the vast majority of tests in the UK and Northern Ireland, and we are not aware of any tests that are routinely sent to laboratories outside the UK. We excluded all results from individuals with the same name and date of birth, referrals from foreign laboratories, and tertiary referrals of previously diagnosed cases to Professor John Newsom-Davis in Oxford. In the 3183 remaining cases (Birmingham 1276, Oxford 989, Sheffield 302, Liverpool 164, St George’s London 112, Glasgow 138, Bristol 100, Cardiff 53, Nottingham 49), we studied age and sex at the time of sampling. Twelve were from babies with neonatal myasthenia gravis and were excluded from further analysis. In 2795 cases (87.8%) an AChR antibody titre was also available.
Estimation of age and sex specific incidence

Given the high specificity of the AChR antibody assay, the fact that the test is rarely requested without clinical suspicion of myasthenia gravis, and that it is routine practice to confirm clinical diagnoses by antibody testing, we considered it reasonable to use the national AChR antibody data to estimate the likely incidence of clinically diagnosed cases of myasthenia gravis. We assumed that repeat testing following initial diagnosis was rare, and that the vast majority of positive titres reported during our three year study period represented incident cases. This assumption was supported by analysis of all tests submitted to the Oxford laboratory from 1994 to 1999; less than 5% of the positive results identified during 1997 to 1999 were from patients diagnosed during 1994 to 1996. Assuming, therefore, that we had an accurate assessment of all positive AChR antibody results in Britain from 1997 to 1999, and that these were incident cases, we used the age and sex stratified mid-year population estimates for England, Wales, Northern Ireland, and Scotland for 1998 to calculate the age and sex specific incidence of AChR antibody positivity in Britain for the period 1997 to 1999. Calculation of 95% confidence intervals (CI) for the incidence rates assumed that the number of cases in each age group followed a Poisson distribution.

Cohort of older people

Blood samples were obtained from the Oxford healthy ageing project, a module of the Medical Research Council cognitive function and aging study (CFAS). Potential participants in the Oxford healthy aging project were nominated as a systematic sample from the primary care registers of all people aged 65 and over, resident in the four central postal districts of Oxford. From a sampling frame of 3555 individuals, 2740 potential participants agreed to be interviewed. The 1649 blood samples were from those who consented to the initial research interview, which included clinical data, and who, two years later, consented to donate a blood sample. The specimens were obtained and stored at −80°C. We also analysed sera from a further 351 patients aged 60 and over attending outpatient clinics with non-neurological complaints at the John Radcliffe Hospital between October 1999 and January 2000. The sera were screened by the immunoprecipitation assay, as used in routine diagnostic testing, and all positive sera were retested to confirm AChR antibody specificity. The Oxford healthy aging project and the associated blood sampling and anonymised analysis were approved by the Central Oxford research ethics committee.

RESULTS

AChR antibodies in referred samples

The age at referral of the 3171 positive AChR antibody results from the UK is shown in fig 1A. The age and sex distribution was consistent across the centres (Birmingham v Oxford v others), and for each of the three years of study (data not shown). The overall incidence was 1.8 cases/100 000 per year. There was the expected high female to male ratio in younger patients, presenting mainly between 15 and 40 years of age, and an increasing number of both men and women presenting...
over 45 years of age, the number of cases peaking between 70 and 80 years and then falling abruptly after age 80 (fig 1A).

In order to calculate the age specific incidence of AChR antibody positivity, the incident data need to be adjusted for the total number of individuals in each age group. In both men and women, the age specific annual incidence (fig 1B) rose sharply between the ages 45 and 74, reaching 4.8 (95% CI, 4.1 to 5.5) per 100 000 in women and 9.9 (95% CI, 8.7 to 11.2) per 100 000 in men at age 75–79 years. However, in both sexes, this trend reversed above age 75–79, and the age specific incidence rates subsequently fell sharply, dropping to 1.6 (95% CI, 0.9 to 2.7) per 100 000 in women and 4.0 (95% CI, 1.9 to 7.4) per 100 000 in men at age 90–94 years (fig 1B).

It was possible that the positive AChR antibody values in older individuals were of less clinical significance than those in younger patients. Figure 2 shows the mean AChR antibody values plotted for different ages at referral for men and women from the Oxford laboratory. Results from other centres were similar (data not shown). Although, as in all studies of AChR antibodies, the range of values is very wide, there was no trend towards lower values with increasing age.

**AChR antibodies in cohort of older people**

Sera from nine of the 2000 individuals aged 60 years and over were positive for AChR antibodies. However, because the results in fig 1 suggested that there might be underreferral of samples after age 75, we separated the samples into those from individuals aged 60–74 and those aged ≥ 75 years. Table 1 shows the age and sex distribution of the 1147 individuals aged ≥ 75 years. Whereas only one of the 853 individuals aged 60–74 years (0.12%) was positive for AChR antibodies, eight of those aged ≥ 75 years were positive (0.71%) (7980 in the community based cohort and 1167 from the outpatient cohort). All but one of the values were at the lower end of the range, but all were compatible with a clinical diagnosis of myasthenia gravis. The AChR antibody values, and clinical data available from the patient records, are summarised in table 2. One individual had had a clinical diagnosis of myasthenia gravis, but no diagnosis of myasthenia gravis had been made in the other seven people with positive AChR antibodies. Notably, however, four of them had a history of stroke or transient ischaemic attacks.

**DISCUSSION**

This national study is the largest ever of the incidence of AChR antibodies. The results confirm previous studies showing a high incidence of diagnosed myasthenia gravis in older people.2–10 In addition, the large size of our study allowed us to determine the age and sex specific incidence of the disease with narrow confidence intervals in the very old. Although the incidence increased steeply in both sexes between the ages of 45 and 79 years, achieving 9.9 per 100 000 in men, it then fell sharply at later ages. This suggests either that myasthenia gravis has a temporary peak in incidence in the older age groups, which then falls off, or that the diagnosis is being missed in many older individuals, particularly in those over 80 years of age. To test our conclusion that there are significant numbers of elderly people with undiagnosed positive AChR antibodies, we studied 2000 individuals aged 60 and over. We found positive AChR antibodies in eight of the 1147 individuals who were ≥ 75 years (prevalence of one in 141), compared with only one in the 853 individuals between 60 and 74 years. Thus the results of both our national incidence study and our community based prevalence study suggest that underdiagnosis of myasthenia gravis may be common in the elderly, and highlight the need for a high index of suspicion in older patients for this treatable disease.

The most striking observation was that the incidence peaked in 75–79 year old men and women and then fell off in people over 80 years old, suggesting underdiagnosis in the very elderly. We could have overestimated the incidence of positive AChR antibody tests in the UK because we could not be certain that all of the tests that we studied were incident cases. However, we took pains to avoid this (Methods), and as we were mainly interested in the age specific pattern rather than the absolute incidence, any overestimation is likely to have applied fairly equally at all ages, and would not explain the steep fall in later life. It could be that we are seeing a “cohort” effect, whereby people born at a certain time, in the early to mid 1920s, had a particular environmental exposure that increased their risk of developing myasthenia gravis. Although a very interesting possibility, the rise and fall in incidence is so steep that it would have required a very short period of exposure, a short period of susceptibility to the exposure (for example, in utero or at the age of 0 to 5 years),

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Demographic characteristics of 1147 individuals aged ≥ 75 years</th>
</tr>
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<tbody>
<tr>
<td>Age (years)</td>
<td>75 to 79</td>
</tr>
<tr>
<td>Men</td>
<td>189</td>
</tr>
<tr>
<td>Women</td>
<td>238</td>
</tr>
<tr>
<td>Total</td>
<td>427</td>
</tr>
<tr>
<td>Positive AChR antibodies</td>
<td>3F</td>
</tr>
</tbody>
</table>

Values are numbers of individuals.

AChR, acetylcholine receptor; F, female; M, male.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Acetylcholine receptor (AChR) antibody positive individuals from among 1147 subjects aged ≥ 75 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, sex</td>
<td>AChR antibody (nM)</td>
</tr>
<tr>
<td>84, F</td>
<td>0.9</td>
</tr>
<tr>
<td>88, F</td>
<td>0.9</td>
</tr>
<tr>
<td>82, F</td>
<td>1.0</td>
</tr>
<tr>
<td>76, F</td>
<td>1.9</td>
</tr>
<tr>
<td>75, F</td>
<td>3.1</td>
</tr>
<tr>
<td>78, F</td>
<td>3.2</td>
</tr>
<tr>
<td>88, F</td>
<td>3.9</td>
</tr>
<tr>
<td>95, M</td>
<td>13.5</td>
</tr>
</tbody>
</table>

F, female; LVF, left ventricular failure; M, male; TIA, transient ischaemic attacks.
and a very long and relatively fixed incubation period (for example, 70 ± 5 years). This seems unlikely given that studies from other countries like Greece have seen a similar peak. Nevertheless, another analysis in, say, 10 years time could be undertaken to address this possibility. Alternatively, the apparent fall off in incidence in the very elderly might reflect competing risks if those who are at risk of developing myasthenia gravis are also at a higher risk than the general population of developing some other, fatal disease. However, for the same reasons given above, the incidence of this disease would have to be extremely strongly associated with the risk of developing myasthenia gravis, and would have to be confined mainly to people in their 80s and 90s.

The overall incidence rate of 1.8/100 000 that we found is higher than the value of 0.74/100 000 in the recent Greek study, which also identified incident cases by a positive AChR antibody test. They found a point prevalence of 17.5/100 000 in individuals aged ≥ 70 years, which is threefold greater than the prevalence of about 5/100 000 in individuals aged ≥ 80 years in a recent UK survey. We have not attempted to calculate the likely prevalence in elderly individuals based on our incidence figures, but one would predict a value higher than either of these studies.

The frequency of positive AChR antibodies in our prevalence study was very surprising. One possible concern is that the high prevalence in individuals aged ≥ 74 years was part of a non-specific increase in autoantibodies with age, and was not of clinical significance. However, this is not supported by the fact that these sera were negative for all other autoantibodies that we examined in parallel (voltage gated calcium and potassium channels, glutamic acid decarboxylase, Hu and Yo antigens) (Clover L, Vincent A, in preparation), and only one of the eight was positive for thyroid antibodies. Moreover, although false positive AChR antibody results can occur in patients on penicillamine treatment, in patients with thymic tumours, and in the monozygotic twin siblings of patients with myasthenia gravis, the assay is otherwise highly specific (> 99.9%) and is thought to be diagnostic of the disease. Further studies are now required to confirm these striking findings in a prospective study, and to investigate their clinical relevance.

Underdiagnosis of myasthenia gravis could be explained at least in part by the fact that many of the presenting symptoms (fatigue, muscle weakness, slurred speech, and difficulty in swallowing) have a broad differential diagnosis in older people. For example, myasthenia gravis is sometimes misdiagnosed as stroke or transient ischaemic attacks (table 2). Unfortunately, under the terms of the ethics approval for the Oxford healthy aging study, we were unable to re-examine any of the AChR antibody positive individuals for overt or subclinical myasthenia gravis. Such a study needs to be done in future cohorts.

There is also evidence of possible underdiagnosis of other uncommon neurological conditions in older people. For example, misdiagnosis and underinvestigation have been suggested as the cause of the apparent sharp fall in incidence of motor neurone disease in late life. Much has been written about the importance of not discriminating against older people in relation to the provision of treatment. It is important also that the same principles are applied to referral for specialist investigation and diagnosis.

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