The presence of glutamic acid at positions 71 or 74 in pocket 4 of the HLA-DRβ1 chain is associated with the clinical course of multiple sclerosis

J M Greer, M P Pender

**Background:** Primary progressive multiple sclerosis (PP-MS) differs from relapsing-remitting or secondary progressive MS (RR/SP-MS) in ways suggesting differences in the pathogenic pathways. Susceptibility to both PP-MS and RR/SP-MS is linked to carriage of the HLA molecule DRB1*1501. Several serologically defined HLA-DR groups (DR1, DR4, DR6, and DR9) occur less often in RR/SP-MS than in controls. Some or all of the HLA-DR molecules encoded by alleles in these serologically defined groups have a negatively charged glutamic acid at residue 71 or 74 of the β1 chain (β171/β174). Residues at these positions are important in the formation of pocket 4 in the antigen binding site of the HLA-DR molecule.

**Objectives:** To investigate whether the presence of alleles encoding HLA-DR molecules containing glutamic acid at β171/β174 correlates with the course of MS.

**Methods:** HLA-DR and HLA-DQ alleles and genotypes were analysed in 121 MS patients (50 with PP-MS) and 109 controls by molecular typing.

**Results:** Alleles encoding HLA-DR molecules containing a glutamic acid at β171/β174 occurred less often in patients with RR/SP-MS than in those with PP-MS or controls. In subjects not carrying the DRB1*1501 allele, a much higher proportion of PP-MS patients carried alleles encoding HLA-DR molecules containing a glutamic acid at β171/β174 than did RR/SP-MS patients or controls.

**Conclusions:** The amino acid residues involved in determining the shape and charge of pocket 4 of the HLA-DR β1 chain could influence the clinical course of MS by determining protection against RR/SP-MS or susceptibility to the development of PP-MS.

Multiple sclerosis (MS) is a chronic inflammatory demyelinating disease of the central nervous system (CNS). Typically, MS has a relapsing–remitting course with clinical attacks followed by some improvement and attack-free intervals. However, after 10–15 years of a relapsing–remitting disease course (RR-MS), more than 50% of patients are likely to have developed a progressive course, characterised by a slow inexorable decline in neurological function. This form of progressive MS is known as secondary progressive MS (SP-MS). In 10–20% of cases of MS, a progressive course occurs from the onset of disease. This form of MS is known as primary progressive MS (PP-MS). PP-MS differs from RR/SP-MS in the amount of CNS inflammation,³ age at onset,³ the female to male ratio,³ the initial symptoms,⁴ and prognosis with regard to disability.⁵ Factors that determine the particular disease course a patient will follow remain obscure, although an older age at onset correlates with a greater likelihood of PP-MS.⁶ Natural history studies, however, show little difference in the progressive phases of PP-MS and SP-MS.⁶ This and the recent report of a patient with subclinical disease for at least 10 years before the clinical onset of PP-MS⁷ are consistent with the suggestion that PP-MS may differ from RR/SP-MS only in having a subclinical initial stage of disease.⁸ Both PP-MS and RR/SP-MS are associated with an increased occurrence of other autoimmune diseases in affected individuals and their first degree relatives, suggesting that they are both autoimmune disorders.⁹ The pathogenesis of MS is thought to involve the development of autoimmunity against components of CNS myelin. This autoimmunity appears to be driven by major histocompatibility complex (MHC) class II restricted T cells,¹⁰ which cannot recognise antigen directly but require that it is presented to them on the surface of antigen presenting cells in association with the MHC class II molecules, which in humans are termed human leucocyte antigen (HLA)-DR, HLA-DQ, or HLA-DP. The HLA molecules are formed from two polypeptide chains, α and β. Some of these chains are extremely polymorphic, particularly the HLA-DR β1 chain and the HLA-DQ α and β chains. The amino acid sequences of the α and β chains of the HLA molecule determine which antigens can be bound and presented to T cells. Immunogenetic studies in MS have shown associations between a limited number of ancestral haplotypes and susceptibility to MS.¹¹ ¹⁴ In particular, the haplotype DRB1*1501–DQA1*0102–DQB1*0602 (that is, the 1501 allele of the gene encoding the HLA-DR β1 chain, the 0102 allele encoding the HLA-DQ α chain, and the 0602 allele encoding the HLA-DQ β chain) correlates with development of MS in northern Europeans, northern European derived populations, Ashkenazi and non-Ashkenazi Jews, Turkish, Mexican Mestizo, and some Japanese and Sardinian patients. However, carriage of these alleles does not correlate with the disease course in MS, as patients with PP-MS and those with RR/SP-MS carry this haplotype in approximately equal proportions.¹⁵ ¹⁷ ¹⁸ ¹⁹ Notwithstanding the lack of correlation between DRB1*1501 haplotype and disease course, the finding that specific immune T cell and antibody responses against myelin and other CNS antigens differ between PP-MS and RR/SP-MS suggests that there may be differences in the other HLA alleles carried by people with different disease courses. However, while some other HLA alleles have been found to be

**Abbreviations:** MS, multiple sclerosis; PP, primary progressive; RR, relapsing-remitting; SP, secondary progressive


Received 25 March 2004
In revised form 13 July 2004
Accepted 9 August 2004

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**Acknowledgments:** We thank all the patients who participated in the study. This work was supported by grants from the National Health and Medical Research Council (NHMRC).
expressed at increased or decreased proportions in either PP-MS or RR/SP-MS compared with controls, no significant differences have consistently been found between PP-MS and RR/SP-MS. Several studies have suggested a link between carriage of DR4 alleles and PP-MS, but others have not. This disparity has largely reflected the relatively small sample size of PP-MS in each study (less than 20 individuals) and the large number of HLA alleles being compared. Recently, however, two studies with larger PP-MS sample sizes have been reported—one from Sweden, which tested 85 PP-MS patients, 727 RR/SP-MS patients, and 455 controls, and one from Ireland, which tested 102 PP-MS patients, 202 RR/SP-MS patients and 398 controls. Interestingly, both studies showed significantly decreased frequencies of DR1, DR4, DR6, and DR9 serologically defined HLA groups in RR/SP-MS patients compared with controls, but no significant differences between PP-MS patients and controls; however, this finding was not specifically mentioned in these papers.

The serologically defined HLA nomenclature (DR1, DR2, DR3, and so on) is often used to report HLA-DR results, as it groups together alleles that show similar reactivity to a panel of tissue typing sera. However, it is now known that each of these sera recognises more than one HLA amino acid sequence. For example, the DR4 serum recognises more than 40 different HLA molecules, each differing slightly in the amino acid sequence of the b1 chain. These slight differences in sequence can greatly affect the ability of a particular HLA molecule to bind and present an antigen. Thus the finding of decreased frequencies of DR1, DR4, DR6, and DR9 in RR/SP-MS patients compared with controls—but no significant differences between PP-MS patients and controls—is of interest because DR1, DR4, DR6, and DR9, together with DR5, are the only serological groupings that contain alleles encoding HLA-DR molecules having a negatively charged glutamic acid at residue 71 or 74 of the b1 chain (b171 or b174).

These two residues—together with residues 13, 70, and 78 of the b1 chain, and residues 9 and 62 of the a chain—are important in determining the shape and charge of pocket 4 of the antigen-binding groove of the HLA-DR molecule.26 27 (Fig 1). However, not all of the alleles in DR1, DR4, or DR6 serological groups encode HLA-DR molecules containing a glutamic acid at b171 or b174. As the HLA data in the two large studies were reported only at the level of serological groups, it is not possible to determine whether the presence of a glutamic acid residue at these positions is correlated with the course of MS. The aim of the current study, therefore, was to test the hypothesis that the frequency of HLA-DRB1 alleles encoding HLA-DR molecules containing glutamic acid at b171 or b174 differ in PP-MS and RR/SP-MS. The results show that such alleles occur significantly less often in RR/SP-MS than in PP-MS, and support the hypothesis that, in some cases at least, HLA can influence the clinical course of MS.

METHODS
Subjects
The 230 individuals used in this study included 50 patients with PP-MS, 71 with RR-MS or SP-MS, and 109 unrelated individuals of white ethnicity as controls. More than 95% of individuals were of northern European descent, with the remainder being of southern European descent. The MS patients were from Queensland and northern New South Wales in Australia and were seen at the MS clinic, Royal Brisbane and Women’s Hospital. All MS patients met the criteria of Poser et al28 for definite MS. All RR/SP-MS patients also met the criteria of McDonald et al19 for MS. Five PP-MS patients did not meet the latter criteria, because magnetic resonance imaging scans were not available for inspection; however, as all these patients had had typical symptoms and signs of PP-MS (with no superimposed relapses) for more than 10 years they were included in the study.

DNA samples
Peripheral blood samples (5–10 ml) were collected by venepuncture into heparinised tubes, after informed written consent had been obtained from each subject. Red cells were lysed and the cell pellet incubated at 37°C overnight in the presence of 0.5 mg/ml proteinase K. High molecular weight DNA was then salted out, using the method of Miller et al.29

HLA typing
HLA-DRB1, -DQA1, and -DQB1 typing was carried out for each DNA sample using Dynal® sequence specific primer sets according to the manufacturer’s instructions. Results of the typing are given as specific alleles, and are also grouped according to serologically defined groups listed by the WHO nomenclature committee for factors of the HLA system.30

Statistical analysis
The frequencies of HLA alleles and genotypes in patients and controls were compared using x2 analysis by 2x2 contingency tables. Yates’s correction was applied when the number of positive samples in any test was less than 5. Fisher’s exact test was used when the total number of samples was less than 50. Odds ratios (OR) and 95% confidence intervals (CI) of the odds ratio for phenotype or genotype frequencies were also calculated. Probability (p) values were corrected for multiple comparisons using Bonferroni’s method, according to the formula: p_corrected (p) = 1 − (1 − p)n, where p is the uncorrected p value and n is the number of comparisons.
RESULTS
HLA-DRB1 phenotype frequencies

The phenotype frequencies of HLA-DRB1 alleles were determined for 50 PP-MS, 71 RR/SP-MS, and 109 non-MS controls. In all, there were 39 different HLA-DRB1 alleles in the MS patients or controls in this study. In order to determine if these populations were similar to the two previous large HLA-DR studies, where results were given as serologically defined groups, the data were analysed and compared with results from those two studies (table 1). In general, the frequencies of serologically defined HLA groups found in our study were within 1–3% of those found previously. The frequencies that we found for DR6 carriage by all individuals differ more; however, they lie within the range of frequencies found in the two large studies (25.9%, 22.7%, and 34.9% for PP-MS, RR/SP-MS, and controls, respectively, in the study from Sweden5; and 14.7%, 9.4%, and 22.7%, and 34.9% for PP-MS, RR/SP-MS, and controls, respectively, in the study from Sweden6; and 14.7%, 9.4%, and 18.3% in the study from Northern Ireland7). The frequencies of DR2 did not differ in non-MS controls or PP-MS in our study and the other two studies.

There were no significant differences between carriage of different serologically defined HLA groups in PP-MS vs RR/SP-MS after correction for multiple comparisons. However, as mentioned in the introduction, an interesting observation was that the difference was much more significant (after p values had been corrected for 19 comparisons) between the total number of RR/SP-MS patients (pooled from the present study and the two previous studies) and total number of controls than between the total number of PP-MS patients and the total number of controls for carriage of DR1 (p = 1.6×10^{-5} v p = 0.999), DR4 (p = 5.0×10^{-5} v p = 0.958), DR6 (p = 4.2×10^{-3} v p = 0.767), and DR9 (p = 6.7×10^{-3} v p = 0.999). This is not just because of the larger numbers of RR/SP-MS patients, and is interesting because a common feature of these serologically defined groups is that they all include alleles that encode HLA-DR molecules containing a glutamic acid at β171 or β174. Indeed, all of the other alleles within these groups that were carried by the individuals studied (both MS and controls) encode proteins that have a positively charged arginine or lysine residue at residue 71 and an alanine at residue 74.

Most other studies reported using molecular typing for HLA either do not show data for all alleles, or use a mixture of PP-MS and RR/SP-MS patients. However, Zipp et al10 reported full HLA-DR typing for a group of 66 patients with RR/SP-MS and 210 controls. In that dataset, there was also a significant decrease in the proportion of RR/SP-MS patients carrying alleles that encode HLA-DR molecules containing a glutamic acid at β171 or β174 (16 of 66) compared with controls (93 of 210) (OR = 0.4 (0.21 to 0.75); p = 0.039), confirming our observations.

HLA-DRB1 genotype analysis

Whereas a small proportion of MS patients are homozygous for their HLA antigens, most express two different alleles at each locus. There is some restriction in the number of different HLA-DRB1 genotypes carried by MS patients compared with controls. This is influenced by the increased frequency of the DRB1*1501 allele in MS; in our study, both RR/SP-MS and PP-MS HLA-DRB1 genotypes contained this allele significantly more often (61.4% and 62.0%, respectively) than control genotypes (23.9%; OR for PP-MS v control = 5.2 (2.5 to 10.7); p = 1.4×10^{-4}, and OR for RR/SP-MS v controls = 8.0 (3.9 to 16.5); p = 1.5×10^{-7}; p values are corrected for 42 comparisons). Of the DRB1 genotypes found in PP-MS, 19 did not occur in RR/SP-MS patients. 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14 of these 19 genotypes contained an allele encoding HLA-DR molecules using a glutamic acid at \(\beta1_{71}\) or \(\beta1_{74}\). In contrast, of 15 genotypes occurring in patients with RR/SP-MS, but not in patients with PP-MS, only four contained an allele encoding HLA-DR molecules with a glutamic acid at \(\beta1_{71}\) or \(\beta1_{74}\).

The percentages of DRB1*1501 positive or negative individuals whose genotypes contained an allele encoding a HLA-DRB1 molecule with a glutamic acid residue at \(\beta1_{71}\) or \(\beta1_{74}\) are shown in Table 3. The percentages of PP-MS patients carrying such alleles were significantly increased compared with the corresponding percentages of RR/SP-MS patients in both the DRB1*1501 positive and DRB1*1501 negative groups. In the DRB1*1501 positive group, this difference was almost entirely caused by DR6 alleles, which made up 78% of alleles encoding HLA-DR molecules with a glutamic acid residue at \(\beta1_{71}\) or \(\beta1_{74}\); however, in the DRB1*1501 negative group, both DR4 and DR6 alleles contributed equally. None of the MS patients had genotypes encoding two HLA-DR molecules that both had glutamic acid residues at \(\beta1_{71}\) or \(\beta1_{74}\), but 5% of controls carried such genotypes.

**HLA-DQA1 and HLA-DQB1 allele genotype frequencies in PP-MS and RR/SP-MS**

In insulin dependent diabetes mellitus, susceptibility to disease has been linked to carriage of particular HLA-DQ alleles rather than to the HLA-DR alleles with which they are in linkage disequilibrium. In MS, it has been suggested that HLA-DQ is more closely associated with disease in Afro-Brazilian patients than is HLA-DR\(^+\); however, in white MS patients, HLA-DR linkages appear to be the strongest. We investigated whether there were differences between PP-MS and RR/SP-MS in carriage of HLA-DQ alleles. The frequencies of individuals carrying different HLA-DQA1 and HLA-DQB1 alleles are shown in Table 4. As expected, DQA1*0102 and DQB1*0602— which are in strong linkage disequilibrium with DRB1*1501— were found significantly more often in PP-MS (\(p_c = 0.03\)) and RR/SP-MS (\(p_c = 0.03\)) compared with controls (\(p = 0.27\) to 0.54; \(p_c = 0.038\)).

### DISCUSSION

The question of whether or not the HLA molecules carried by a patient with MS can influence the clinical course of disease
typing was presented only in terms of the serological groups. Thus it is not possible to estimate from those studies the percentage of PP-MS patients carrying alleles containing a negatively charged glutamic acid residue in pocket 4. There are no other published molecular typing studies using well defined PP-MS patients.

The amino acids found at β171 or β174 of the HLA molecule, together with those at positions 13, 70, and 78 of the β1 chain, and at positions 9 and 62 of the α chain, influence the shape and charge of pocket 4 of the particular HLA-DR molecule.27 This is illustrated in fig 1 for DRB1*0404 and DRB1*0407, which differ in pocket 4 only at β171; this residue is an alanine (non-charged, hydrophobic) in DRB1*0404 and a glutamic acid (acidic side chain) in DRB1*0407.

Pocket 4 has been shown to be critically important in the binding of the immunodominant peptide (residues 84–102) of myelin basic protein, a putative autoantigen in MS,10 to DRB1*1501 in patients with RR-MS.44 For that particular peptide/HLA combination, a large, predominantly hydrophobic uncharged pocket 4 is required to accommodate the large aromatic side chain of a phenylalanine residue. Studies in other MHC class II restricted autoimmune diseases such as rheumatoid arthritis, pemphigus vulgaris, systemic lupus erythematosus, and Hashimoto’s thyroiditis have also shown that β171 and/or β174 play a critical role in the recognition of the HLA-DR/peptide complex by CD4+ helper T cells.45,46 In rheumatoid arthritis, for example, the peptides that bind to proteins encoded by DRB1*04 alleles that are positively (DRB1*0404) or negatively (DRB1*0402) associated with disease have been characterised.47 These peptides all have identical requirements at the anchor residues at peptide positions p1 and p6, but substantially different requirements at position p4 (the residue that binds in pocket 4 of the HLA molecule). DRB1*0402, which has a negatively charged glutamic acid at β171, binds peptides with a positive charge at p4, whereas the presence of the positively charged arginine residue at β171 in DRB1*0404 correlates with binding of peptides with a negative charge at p4.

The pocket 4 region of HLA-DR molecules is also important in non-autoimmune inflammatory diseases mediated by CD4+ helper T cells. For example, the presence of a positively charged arginine at positions β130 or β131 is associated with susceptibility to tuberculosis leprosy.48 In contrast, chronic beryllium disease—a lung disease caused by the accumulation in the lung of beryllium specific CD4+ MHC class II restricted T cells—is associated with the presence of a glutamic acid at residue 69 of the DPB1 chain.49 Residue 69 of the DP β chain corresponds to β171 of the DR chain.

Some studies have suggested a trend towards increased carriage of DR4 in patients with PP-MS compared with those with RR/SP-MS,5,21 although this difference has not reached statistical significance in any study. In the current study, there were no differences in the overall frequency of DR4 alleles in PP-MS compared with RR/SP-MS, although more PP-MS patients carried DRB1*0402 or *0407—which encode HLA-DR molecules containing a glutamic acid residue at β171 or β174—than did RR/SP-MS patients (12.0% vs 4.4%, respectively). In particular, DRB1*1501 negative PP-MS patients showed an increased frequency of these alleles in the genotype compared with DRB1*1501 negative RR/SP-MS patients (p = 0.036 by Fisher’s exact test). It is therefore possible that the previously reported DR4 association with PP-MS5,21 may have reflected the presence of alleles encoding HLA-DR molecules containing glutamic acid at β171 or β174.

The most obvious effect of the change in shape, size, or charge of pocket 4 would be on the ability of the particular HLA molecule to bind a disease relevant peptide; however, it

## Table 4: Phenotype frequencies for HLA-DQA1 and DQB1 alleles

<table>
<thead>
<tr>
<th>DQA1 allele</th>
<th>PP-MS (n = 30)</th>
<th>RR/SP-MS (n = 71)</th>
<th>Controls (n = 109)</th>
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<tr>
<td>0101</td>
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<td>0102</td>
<td>33 (66%)*</td>
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<td>3 (6%)</td>
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<td>0201</td>
<td>7 (14%)</td>
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<tr>
<td>0601</td>
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<td>1 (1.4%)</td>
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<tr>
<th>DQB1 serological group</th>
<th>PP-MS (n = 30)</th>
<th>RR/SP-MS (n = 71)</th>
<th>Controls (n = 109)</th>
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<tr>
<td>DQ5(8)</td>
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<td>13 (18.3%)</td>
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<td>DQ6(1)</td>
<td>38 (76%)*</td>
<td>52 (73.2%)*</td>
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<td>27 (54%)</td>
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<tr>
<td>DQ4</td>
<td>1 (2%)</td>
<td>4 (5.6%)</td>
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*p < 0.03 v controls; *p < 0.002 v controls.

MS, multiple sclerosis; PP, primary progressive; RR, relapsing-remitting; SP, secondary progressive.

has been debated now for almost 20 years. Several factors have made it difficult to draw firm conclusions. These have included unclear definition of patient subgroups (such as the grouping of SP-MS and PP-MS), small PP-MS patient numbers, and the analysis of HLA only at the level of serologically defined groups. Most attention has focused on HLA-DRB1*1501, as this allele correlates with susceptibility to disease in numerous studies.11–13 15–20 21–27–29 However, in HLA heterozygous individuals, who make up the majority of people with MS, the immune system recognises antigen presented not only by DRB1*1501 but also by the HLA molecules encoded by the other alleles in the genotype. In several other autoimmune diseases—including insulin dependent diabetes mellitus, rheumatoid arthritis, and Addison’s disease—it has been shown that combinations of certain alleles in the genotype can affect the development of disease50–52 or the disease course.42–43

Our results suggest that the presence of alleles encoding HLA-DR molecules containing a negatively charged glutamic acid residue at β171 or β174 protects against the development of a relapsing-remitting course or increases the susceptibility to a primary progressive course of disease, or both. HLA-DR genotypes consisting of one of these alleles, with or without DRB1*1501, are significantly decreased in patients with RR/SP-MS in comparison with those with PP-MS. As the antigens involved in the development of the different clinical forms of MS have not been defined, it is difficult to determine whether these alleles actively participate in the development of a primary progressive course. If, as has been suggested,4 PP-MS differs from RR/SP-MS only in that the relapsing-remitting phase is subclinical, then another possible interpretation of the data in the current study is that carriage of alleles encoding HLA-DR molecules containing a negatively charged glutamic acid residue at β171 or β174 prevents the initial relapsing-remitting phase of disease.

This correlation between the clinical course of MS and the occurrence of alleles encoding HLA-DR molecules containing a negatively charged glutamic acid residue in pocket 4 of the antigen binding groove has not been noted previously because only two studies have been published using a group of more than 20 PP-MS patients,5 22 and for both of these HLA...
is also possible that the DR4, DR6, and other HLA molecules containing a glutamic acid at β171 or β174 are encoded by alleles that are in linkage disequilibrium with some other site within the MHC region which could affect the disease course.

We did not find any evidence to suggest that HLA-DQ alleles account for the differences seen between PP-MS and RR/SP-MS (table 4); however, the HLA region is also in strong linkage disequilibrium with genes encoding, for example, the cytokine tumour necrosis factor (TNF) and the molecule IkBα, a member of the IkB family of inhibitory factors that are involved in control of the pro-inflammatory transcription factor, NF-κB. Several polymorphisms are known to occur in TNF and IkBα genes. Patients carrying HLA alleles in linkage disequilibrium with such polymorphisms could have altered inflammatory immune responses, thus leading to a different clinical course of disease.

Not all PP-MS patients carried alleles encoding HLA-DR molecules containing a glutamic acid residue at β171 or β174, and some RR/SP patients did carry such alleles. This probably reflects the heterogeneity in disease pathogenesis. Overall, however, there was a clear cut difference between the two groups of MS patients, which suggests that—in some cases at least—the HLA genotype can exert an influence on the clinical course of MS. The results from the study of Zipp et al.12 and from the large serotyping studies from Sweden6 and Northern Ireland13 provide indirect support for this hypothesis.

ACKNOWLEDGEMENTS

We would like to thank Lynn Mallard, Diane Muller, and Peter Cottrell for excellent technical assistance. This work was supported by grants from the National Multiple Sclerosis Society (RG 3190-A-1), and MS Australia.

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Competing interests: none declared

REFERENCES


Robert Bárány 1876–1936

Robert Bárány was educated at the University of Vienna, graduating in medicine in 1900. In 1903, he accepted an appointment at the University of Vienna otology clinic working in the department of Adam Politzer, who had established the specialty in Austria. Here he began his studies exploring the equilibrium system of the ear and the brain. Among his contributions was a test, now called the Bárány test, for diagnosing diseases of the semicircular canals of the inner ear by syringing the ear with either hot or cold water. He noted nystagmus when the water was above or below, but not at body temperature. He concluded that with caloric nystagmus, the endolymph, normally at body temperature, displaced itself when it came into contact with water at other temperatures, and that the vestibular nerve transmitted the thermal disturbance within the endolymph to the brain. His work led to the development of the Bárány rotating chair, which demonstrated the relationship between the vestibular system, the nervous system, and centres within the cerebellum. In 1913, he investigated the somatopic localisation in the cerebellum and postulated that the vermis was concerned with coordinated movements of the trunk, the hemisphere with the extremities, and the flocculus with movements of the eyes.

Bárány received the Nobel Prize in 1914 for his development of clinical investigation of the human equilibrium system. He essentially founded a new area of medical science and practice. At the time of this award, which was delayed until 1915 by the outbreak of World War I, he was a prisoner of war in Siberia. When Austria entered the war, Bárány saw an opportunity to investigate his ideas with brain injured soldiers. He volunteered for medical service and was assigned to the fortress of Przemysl in Galicia (now Poland) to provide care for 125,000 men. There he built an otolaryngology unit, and developed a very successful primary suture technique for the suture of head wounds. He was captured there by advancing Russian troops in April 1915 and taken prisoner. After Prince Carl of Sweden persuaded Czar Nicholas I to release Bárány, the Swedish Red Cross negotiated his release and emigration to Sweden. He became director of the otorhinolaryngology clinic at the University of Uppsala, Sweden in 1917, and in 1926 became a full professor there. In 1921 he became the first to describe positional vertigo.

Upon his death, the University of Uppsala established the Bárány medal, to be awarded at 5 year intervals to the most outstanding scientist in the field of the vestibular research. The Bárány society was founded in his honour by C S Hallpike and C O Nylen in 1960.

Bárány is portrayed here on a stamp issued by Austria in 1976, the centenary of his birth (Stanley Gibbons no. 1756, Scott no. 1031).