Sib risk and the dizygotic twin concordance rate for multiple sclerosis

WILLIAM H JAMES

From the MRC Mammalian Development Unit, Wolfson House (University College London), 4 Stephenson Way, London NW1 2HE

SUMMARY Data have been reviewed on sib risk and the dizygotic twin concordance rate in multiple sclerosis. Even when rigorous criteria are applied, the dizygotic twin concordance rate for multiple sclerosis is apparently higher (perhaps 10 times higher) than could be explained by the sib risk. In contrast, twins with Parkinson's disease have low concordance rates even when ascertainment is by informal methods. It is concluded that such methods of ascertainment are not as biased as has been suggested, and that the high concordance rates reported for multiple sclerosis are a characteristic of the disease rather than an artifact of the ascertainment. Three hypotheses are considered which might, in principle, explain this high dizygotic twin concordance rate in multiple sclerosis:

1. One is certainly false, viz, that it is due to an excessive liability of dizygotic twins to the disease.
2. It is possible that a pathogen occurs in early infancy or in pregnancy itself.
3. It seems more likely that the high concordance rate may be explained in terms of age related events or sequences of events. (If such events were pathogenic for one member of a sibship, they would be pathogenic for another only if it were a co-twin.)

I suggested\(^1\) that the concordance rate for multiple sclerosis (MS) in dizygotic (DZ) twins may be higher than can be accounted for by the sib risk; and a similar suggestion has since been made by other workers.\(^2\) However, as the suggestion has met with some scepticism,\(^3\) I want to review the evidence on both sorts of data involved in that contrast. These data on MS will also be compared with analogous data on Parkinson's disease.

A review of data on sib risk and DZ twin concordance for MS

Tables 1 and 2 summarise all the relevant empirical data on MS known to me.

If concordance in twins were due merely to the known raised sib risk, then \( r = 2c \), where \( r \) is the sib risk and \( c \) the concordance rate (proportion concordant among affected twin pairs).

It is perfectly clear that if these data were accepted at face value, then the concordance rate is far higher than can be accounted for by the sib risk. The sib risk seems to be of the order of 1\(^{\%}\), and the concordance rate of the order of 10\(^{\%}\), a value which prima facie is 20 times too high for the null hypothesis.

Accordingly, the data must be scrutinised. It is a matter of judgment to decide to what extent they may be accepted. To facilitate this judgment, let us consider the possible sources of bias. Interest here attaches to biases occasioning high estimates of concordance rates or low estimates of sib risks. A catalogue of the various sorts of bias is now offered with comments on their relevance to this particular interest.

The biases

1. Multiple sclerosis is a difficult disease to diagnose; in many cases firm diagnosis is not made until necropsy. The result of this uncertainty is that living patients are usually classified as 'probable' or 'possible'. This leads to a range of categories (rather than 'concordant' or 'discordant') into which pairs of sibs may be classified. However none of these considerations seems to suggest that estimates of concordance rates are more biased than, or biased in a different direction from, those of sib risks.

2. Sib risks and twin concordance rates are both (roughly equally) underestimated because a sib or twin, who is unaffected at the time of survey, may later develop the disease. It is true that if the average
Table 1 Numbers of DZ twin pairs concordant and discordant for MS reported by various authors

<table>
<thead>
<tr>
<th>Author</th>
<th>Concordant pairs</th>
<th>Discordant pairs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heitberg &amp; Holm</td>
<td>1</td>
<td>27</td>
</tr>
<tr>
<td>Bobowick et al</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>Currier &amp; Eldridge</td>
<td>3</td>
<td>26</td>
</tr>
<tr>
<td>Mackay &amp; Myrianthopoulos</td>
<td>6</td>
<td>23</td>
</tr>
<tr>
<td>Schapira et al</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Thoms</td>
<td>17</td>
<td>29</td>
</tr>
<tr>
<td>Koch</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Bammer et al</td>
<td>1 + 1</td>
<td>5</td>
</tr>
<tr>
<td>Cendrowski</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Schwermans</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>Williams et al</td>
<td>2</td>
<td>10</td>
</tr>
</tbody>
</table>

See notes on p. 42

Table 2 Numbers of cases of MS in sibs of index cases

<table>
<thead>
<tr>
<th>Place</th>
<th>Total</th>
<th>Affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Switzerland</td>
<td>Curtius &amp; Speer</td>
<td>444</td>
</tr>
<tr>
<td>England &amp; Wales</td>
<td>Pratt et al</td>
<td>538</td>
</tr>
<tr>
<td>Sweden</td>
<td>Muller</td>
<td>2268</td>
</tr>
<tr>
<td>Northern Ireland</td>
<td>Allison &amp; Millar</td>
<td>2939</td>
</tr>
<tr>
<td>Northern Scotland</td>
<td>Sutherland</td>
<td>547</td>
</tr>
<tr>
<td>Northumberland &amp; Durham</td>
<td>Schapira et al</td>
<td>2151</td>
</tr>
<tr>
<td>Vestfold, Norway</td>
<td>Oftedal</td>
<td>554</td>
</tr>
<tr>
<td>Møre og Romsdal, Norway</td>
<td>Presthus</td>
<td>335</td>
</tr>
<tr>
<td>British Columbia</td>
<td>Sadovnick &amp; MacLeod</td>
<td>1179</td>
</tr>
<tr>
<td>United States</td>
<td>Mackay &amp; Myrianthopoulos</td>
<td>207</td>
</tr>
<tr>
<td></td>
<td>∑</td>
<td>11162</td>
</tr>
</tbody>
</table>

By pooling all these data, one may derive a sib risk of 149/11 162 = 0.013.
See notes on p. 42

interval between the onset of MS in non-twin sibs were greater than that relating to concordant twins, then some slight underestimate of sib risk (contrasted with twin concordance rate) would result. But it is not clear that the one interval is greater than the other.

3 Both sib risks and twin concordance rates have probably been underestimated (roughly to the same degree) because familial involvement will naturally suggest another diagnosis (eg, spastic ataxia and paraplegia) rather than MS.

4 It is important that pairs of cases (whether of sibs or twins) should be ascertained within series rather than presented on account of their rarity. If this principle is not adhered to, then it is impossible to make a numerical assessment of this rarity. Of all the sources of bias, this alone seems to be of greater magnitude in regard to twins than to other sibs. The sib risks offered in table 2 may be regarded as unbiased estimates at the time of survey. It is not clear that concordant pairs of sibs are more likely to be reported than discordant pairs. However the twin data presented in table 1 cannot all be regarded as unselected. In particular, the data of Mackay and Myrianthopoulos were collected as a result of an appeal for twins with one or both affected. So there is a suspicion that concordant pairs would be more likely to be ascertained.

Analysis of the data on DZ twin concordance in MS
Using the most rigorous criteria (accepting only the data recommended in Note a to table 1) it will be seen that one probable plus two certain concordant pairs occurred among 44 DZ twin pairs affected by MS. If we accept a sib risk of 0.013 (derived from the data in table 2), then there were (44/2) × 0.013 = 0.286 concordant pairs expected. The Poisson probability of 2 or more events occurring when 0.286 are expected is close to 0.03: and the probability of 3 or more occurring is less than 0.005. So there can be no reasonable doubt that more DZ pairs are concordant for MS than would be predicted on the basis of sib risk. The magnitude of this risk factor is of the order of (2 + 17)/0.286 = 10. To offer confidence limits for this factor would attribute a spurious precision to it. However, bearing in mind all the data in table 1, and the notes on them, it still seems that 10 is a reasonable estimate of that factor.

A comparison of MS with Parkinson’s disease
It seems worth contrasting this high DZ twin concordance rate for MS with the comparable figure for Parkinson’s disease (PD). Like MS, this is a progressive neurological disorder of unknown cause. Like MS sufferers, PD patients have a much reduced life expectancy. Both diseases have a very variable age at onset, though the mean age at onset of PD is greater than that at onset of MS. Both diseases present difficulties in diagnosis, each being occasionally misdiagnosed in patients with the other disorder. Both diseases show unequivocal signs at necropsy, but no objective markers are available whereby diagnosis can reliably be made during life. Of interest in the present context is the fact that PD, like MS, has been thought to have some (perhaps weak) genetic determinants. Family histories have been reported to contain PD patients in up to four generations.

The purpose of the present excursus may now be apparent. We are interested in concordance rates in PD twins ascertained with the same sorts of bias as those characterising the ascertainment of some of the MS twins reviewed above. A recent study offers such data. I quote the description of the ascertainment: “Twin pairs in whom one or both were said to have PD were sought through adult twin registries in the United States and through notices in medical journals, notices in newsletters of voluntary PD organisations, announcements at neurology meetings, and informal inquiries among professional
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The results were as follows: among 43 monozygotic (MZ) and 19 DZ pairs in which an index case had definite PD, only one MZ pair was definitely concordant. When pairs with questionable clinical features were included, 4 out of 48 MZ and 1 of 19 DZ pairs were concordant. The authors conclude that: “the concordance rate for MZ twins is low and likely to remain so during extended follow up. The frequency of PD in the MZ co-twins of affected index cases is probably identical to the frequency to be expected in a group of unrelated controls matched for age and sex”.

I suggest that this conclusion (based on the ascertainment described above) indicates that ascertainment by such relatively informal methods does not in itself greatly affect the probability of inclusion of concordant pairs. (The alternative inference is that PD in one member of a twin pair protects the other member from the disease. This view is supported by the fact that among sporadic reports of PD twins, discordant pairs outnumber concordant pairs by 5 to 1.3,7–9

Accordingly, I suggest that the high concordance rates for MS, reported in both MZ and DZ twin pairs, are not wholly artifacts of admittedly imperfect methods of ascertainment but are real features of the disease.

What can be the cause of this? Three hypotheses will be discussed.

HYPOTHESIS 1
That DZ twins are particularly prone to MS
If DZ twins were particularly prone to MS, then DZ twin concordance rates would be expected to be high compared with sib risk. However the facts do not suggest that DZ twins are at additional risk. The only data I have on the incidence of twins in samples of MS patients point decisively in the opposite direction. Cendrowski13, Shapira et al., and Allison & Millar22 together report nine discordant twin pairs among 1635 MS patients. At the time of birth of these patients about 1 in 80 were twins, so about 1 birth in 40 was of a twin. Perinatal mortality and infant mortality are substantially greater in twins, but after the first year the mortality differential seems largely to disappear. So, using the data cited by Bulmer,28 one would expect about 1 twin in 46 adults. Whence one might have expected about 35 instead of the nine twin pairs actually observed in the pooled data cited above. Confirmatory evidence on this point is provided by the data of Bobowick et al.,16 who estimated that among their sample of 16 000 adult male twins aged 53–63, the proportion suffering from MS was about half the proportion estimated on population age specific rates. (In parenthesis it might be remarked that the deficit of MS in twins may be explained by the facts that (a) birth order is positively associated with (dizygotic) twinning, and (b) birth order seems to be negatively associated with MS.23) In conclusion, it is clear that the high concordance rate in DZ twins is not accounted for by an excess of MS in DZ twins.

HYPOTHESIS 2
That a pathogen occurs in pregnancy or infancy
So it seems that DZ twins must somehow experience a greater similarity of environmental exposure than do other pairs of sibs. But how? Several kinds of hypothesis may be formulated. Twins obviously spend more time in one another’s company in infancy (and later) than do other pairs of sibs. This could scarcely account for a 10-fold risk ratio unless a pathogen occurs in infancy or in pregnancy itself. As far as I know, little attention has been paid to this possibility. It might be worth looking at the in utero experience of patients.

Did the mothers suffer from any dietary deficiency or infection during pregnancy? A point against this hypothesis is that MS seems not to be associated with lower social class.33 Moreover, migration studies have made it fairly clear that a pathogen occurs during childhood,33 though this does not rule out that it (or another pathogen) should sometimes occur earlier.

HYPOTHESIS 3
That MS follows a sequence of events
Suppose the pathogen consists of a sequence of events, each of which must occur during a limited age range (eg, event A at age 2 and event B at age 4). If this were so, and if events A and B occurred in a household at times that were pathogenic for one sib, they might not be so for another unless it was a twin. The point may be illustrated as follows. Suppose (as has been proposed) that MS is dependent on an infection which is harmless in early childhood but pathogenic if it first occurs later in life. (Such a suggestion is supported by the indications that MS is more likely to occur in first born children.31) In such circumstances, one would expect a high concordance rate in DZ twins compared with sib risk.

The hypothesis may be elaborated as follows. It seems to be well established that (in some samples) measles antibody titres are higher in MS patients than in their controls.34 Moreover, the geographical distribution of measles is unlike that of MS; hence the inference that if measles does play some part, then a cofactor is also responsible. Let us, for example, suppose that event B is a measles infection. (This is consistent with the suggestion that MS patients have measles later than controls.33 36 What then about the identity of event A? We need some agent which
increases in prevalence with latitude. One might suggest the common cold, influenza or pneumonia. (These are merely suggested as examples; readers may supply their own.) Of these, pneumonia would be the most attractive to test because (a) it is rarer and would be better recorded, and (b) any hypothesis invoking a very common disease carries the additional burden of explaining why it is not always pathogenic.

I am grateful to Mrs June Rathbone of Galton Laboratory for help with the translation.

Notes to Table 1
These notes deal with the adequacy of sampling of twin pairs affected by MS.

(a) The data of Heltberg & Holm, Bobowick et al, Schapira et al, Bammer et al and Cendrowski may be accepted as adequately ascertained within series.

(b) As noted above, the data of Mackay and Myrianthopoulos are clearly suspect; and, to a lesser extent, so are those of Currier and Eldridge. (Some cases are common to these two sets of data.) The data of Currier and Eldridge were collected via a notice in the patient bulletin of the Multiple Sclerosis Society. Probably less bias attaches to such a proceeding than to a public appeal because patients, administrative workers, and ancillary workers are all likely to be responsive to the research imperative in such a milieu—so presumably almost all twins known to the Society would get ascertained. It seems, however, that any rigorous assessment of the data would dictate exclusion of this material.

(c) It is difficult to know how to deal with the data of Thums. He regarded his single concordant pair as doubtful. It would seem unduly cautious to exclude this pair but to include the discordant pairs; but it might be incautious to include both. So all his material will be excluded here.

(d) The data of Koch are a collection of single cases published separately by different authors. They were not ascertained in series so they must be excluded here.

(e) The series of Schwermann was ascertained from the records of a neurological clinic in Munster. If a patient was identified in these records as a twin, then he/she was included. It is not clear if the recording of the twinship was made as a result of a question which was routinely asked of all patients. If it was, then the ascertainment seems to have been unbiased. If, on the other hand, it depended on the patient volunteering the information, then biased ascertainment may have resulted as a consequence of the fact that a patient would be more likely to volunteer information about a concordant than a discordant co-twin. In face of this uncertainty, these data will be ignored.

(f) The cases of Williams et al were ascertained via a notice in the MS Messenger, the newsletter of the MS Society of the USA. Only like-sexed pairs were accepted. This method of ascertainment is like that of Currier & Eldridge (and one may wonder whether the two series contain pairs in common). So these data cannot be regarded as completely representative.

Notes to Table 2
1 Where some discretion may be applied to the figures to be accepted, I have preferred to risk overestimating (rather than underestimating) the sib risk in order that the contrast with twin concordance rates may be conservative.

2 The data of Hyllested have been omitted because Acheson suggested that they underestimated the sib risk.

3 Muller noted that in the data of Curtius & Speer the diagnosis was not conclusive in any of the four affected sibs: he noted that in two, the diagnosis was ‘even extremely questionable’.

4 The data of Pratt et al contain an affected sib who was dead by the time of the survey.

5 Some uncertainty relates to the data of Oftedal and of Presthus. These authors both seem to suggest that ascertainment was practically complete. But their estimations of sib risk may be appropriate instead to single ascertainment (thus halving the estimate of sib risk). Their estimates were accepted by Berry, but they have been doubled here in order, as argued above, to keep errors conservative. Numerically these were the two smallest studies, so the point is not of practical importance.

6 The cases of Mackay and Myrianthopoulos were ascertained by an affected twin. These authors found nine definite and five possible cases among the 236 sibs of affected twins. However among these 236 sibs, Mackay and Myrianthopoulos included the co-twins of DZ twin propositi. This clearly is an inappropriate method of estimation of sib risk if it is suspected that the DZ twin concordance rate may be high. So the 29 co-twins (including six affected) of DZ twin propositi have been subtracted. This leaves eight definite and possible cases among the 207 non-twin sibs of twin propositi. There are several reasons for expecting this proportion to be higher than other estimates of sib risk:

(a) Mackay & Myrianthopoulos make it clear that these sibships had been followed for an unusually long time, thus allowing the sibs more time to have developed the disease.

(b) Twinning (or at any rate DZ twinning) is associated with positive maternal age and parity effects. So the twin propositi would be expected on average to occur later in their sibships than would non-twin propositi. Ipso facto the sibs of twin propositi would occur earlier. This constitutes an additional reason for supposing that those sibs had an unusually long time to develop the disease.

(c) Lastly I have reviewed the data and suggested that within sibships early born members are more likely to develop MS than later born members (results to the contrary all being based on invalid tests). This being so (and bearing in mind, as argued above, that
the sibs at risk are in this case likely to be early born), one would expect an unusually high proportion of them to be affected.

References

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W H James

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