FAMILIAL INCIDENCE OF DISSEMINATED SCLEROSIS IN NORTHERN IRELAND

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IN Northern Ireland we have recently completed a survey of disseminated sclerosis; 700 cases have been traced and examined and in 44 families we have found more than one member affected, giving a familial incidence of 6.58 per cent.; this figure corresponds closely with the incidence in recent surveys from the Middlesex Hospital 6.5 per cent. (Pratt, et al., 1951), and from the Bristol area 6 per cent. (Campbell, 1952).

It is only in the past decade that the familial aspect of the disease has been generally recognized in the English literature, although the subject was much discussed on the Continent. As recently as 1930 Russell Brain, in a review of the literature of disseminated sclerosis, stated: "In striking contrast to diseases attributable to an inherited germinal defect, multiple cases of disseminated sclerosis in one family are extremely rare compared with sporadic cases, and its occurrence in more than two members of a family and in two successive generations is almost unknown. These facts suggest that inherited predisposition plays no part in the ætiology of the disease, and that the occasional occurrence of multiple cases in one family is due either to chance, exposure to a common environment or mutual infection." Curtius (1933) made an extensive study of the 2.778 near and distant relatives of 56 cases of disseminated sclerosis in Bonn and a less extensive study of 346 relatives of a further 50 cases in Heidelberg. In the Bonn series he found 6 definite cases and in the Heidelberg series 4 definite cases of disseminated sclerosis among the relatives. Later, in 1937, with Speer, he described 2 further families with multiple cases. They found one doubtful case in the 212 parents and 4 definite, and 1 doubtful case in the 444 siblings; 4 in 444 is equivalent to 90 per 10,000 and 40 times the incidence in the general population, based on the Swiss surveys (Bing and Reese, 1926; Ackermann, 1931). As a control group, Curtius investigated the 640 relatives of 56 patients with fractures and found no case of disseminated sclerosis. Mackay (1950) reviewed the literature and, after careful consideration of the case reports, accepted 79 families with multiple cases of disseminated sclerosis. He added a further 5 families. He also found that up to and including 1948, autopsy confirmed the diagnosis in 3 patients in one family, in both patients in 4 families, and in one of two patients in 13 families. Pratt, et al. (1951), found 184 families in the literature where more than one member was affected with disseminated sclerosis. In their series of 310 cases the familial incidence was 6.5 per cent. (20 families). The incidence of disseminated sclerosis in the siblings of 168 cases and the parents of 310 cases was significantly higher than that expected on the basis of a random distribution of the disease.

MATERIAL.

We have attempted to trace all cases of disseminated sclerosis in Northern Ireland. In addition to many patients who have attended the Neurological Clinic, all general practitioners were asked to notify us of the names of patients suffering from the disease. Seven hundred cases were seen, the majority in their homes. This had the advantage that relatives could supply missing details in the clinical and family histories. Frequently it was possible to obtain details concerning the health of grandparents and distant relatives, and in a relatively compact and self-contained community such as Northern Ireland the news that we had visited a relative increased the likelihood of obtaining a positive family history. In 1947 one of us (Millar, 1949) made a limited survey of the disease and traced 91 cases. There were multiple cases in 3 out of 89 families. Some of these 91 cases are included in this survey and two additional familial cases have been found, now making an incidence of 5 in 89 families. This suggests that the greater the scope

DISTRIBUTION	of 700	CASES	IN THE	DIAGNOSTIC	GROUPS
Probable	-	-	-		476
Possible	-	-	-		145
Early -	-	-	-		79

of the study the greater the chance of finding multiple cases in one family. We cannot, of course, claim to have seen every person suffering from this disease in Northern Ireland, but we think we have seen the majority, for reasons given in the first paper (Allison and Millar, 1954). We have been strict in the criteria of diagnosis and cases were placed in the following three groups (Table 1). The criteria for this grouping are discussed in greater detail elsewhere (Allison and Millar, 1954), but briefly "probable" cases are those which show typical evidence of dissemination of lesions with or without a history of remissions. "Possible" cases are those where the findings suggest the diagnosis and no other cause has been found for the clinical picture. "Early" cases are those where the history is suggestive, but where there are few or no neurological signs.

There were 44 families with two or more members affected out of a total of 668 families, giving an incidence of 6.58 per cent. Table 2 shows the relationships. We examined 64 cases in the series and an additional 6 cases not included in the series—70 cases in all. In this total were 56 probable, 11 possible, and 3 early cases. The familial group is again sub-divided into two categories (see Appendices XY for case histories and pedigrees) :—

1. Families in which we had examined two or more members (23 families) or where we had examined one member and the evidence from another centre was sufficiently strong to warrant a firm diagnosis in the other (6 families).

2. Families in which we had examined one member and the evidence concerning the other was less convincing, such as letters from the family doctor, or death certificates (15 families).

In addition, in the Appendix Z we have recorded the incidence of other neurological conditions found in near relatives of our cases. Under the heading

TABLE 2.

Relations	SHIPS			
(including cases dead	or not	t examin	ned).	
Grandfather, father and son	-	-	-	1
Father and son	-	-	-	5
Mother and daughter -	-	-	-	1
Mother and two daughters	-	-	-	1
Mother and son	-	-	-	1
Mother, son and daughter	-	-	-	1
Brother and sister -	-	-	-	13
Brother, sister and her daug	hter	-	-	1
Two brothers and one sister	•	-	-	3
Three brothers	-	-	-	1
Two sisters	-	-	-	8
Two sisters and one brother	-	-	-	1
Uncle and nephew -	-	-	-	2
Aunt and niece	-	-	-	2
Cousins (first)	-	-	-	3
. ,				
Total number of familie	s with	h more	than	
one member affected	-	-	-	44 (6.58%

chronic neurological disorders are included 18 persons about whom remarks such as the following were made :----

"Paralysis of the spine many years before death," "both legs affected, can't walk," "similar complaint to mine." This group may well include further cases of disseminated sclerosis. This is one indication that our estimate of the familial incidence in this series is conservative.

DIFFERENTIAL DIAGNOSIS FROM FAMILIAL NEUROLOGICAL DISORDERS.

Hereditary Spastic Ataxias: This group includes Friedreich's Ataxia (1863), and here the absence of the deep tendon reflexes makes confusion unlikely, although 6 cases were notified in this series. However, regarding the other two components of this group, hereditary ataxia and spastic paraplegia, the differential diagnosis may be very difficult if not impossible on clinical grounds. There are, however, certain points which would be in favour of these two conditions as compared with

TABLE 3.

DISTRIBUTION OF SYMPTOMS AND SIGNS IN THE THREE GROUPS. Expressed in percentages.

NUMBER OF CASES			Fami	lial D.S.	* Proba	ble D.S.	Hered	itary Ataxia
			61	cases	423	cases		18
Remissions	-	-	-	61%		76%	•••	0%
Onset under 20 yea	rs	-	-	11		20		39
Deformities	-	-	-					33
Diplopia -	-	-	-	30		37		22
Retrobulbar neuritis	-	-	-	28		36		_
Paræsthesiæ	-	-	-	45		65	• • • •	22
Urinary symptoms	-	-	-	46		66		28
Weakness of legs	-	-	-	96		92		78
Weakness of arms	-	-	-	33		49		6
Ataxia of legs	-	-	-	33		42		67
Ataxia of arms	-	-	-	35		42		60
Nystagmus -	-	-	-	58		56		44
Monocular nystagm	us	-	-	25		9		
Pale discs with visu	al impai	rment	-	18		27		22
Pale discs without v	isual im	pairme	ent	42		34		2 2
Euphoria -	-	-	-	35		38		28
Dementia -	-	-	-	17		11	• • •	
Dysarthria -	-	-	-	13		25		22
Impaired vibration s	sense	-	-	51		60		28
Impaired postural se	ense	-	-	37		43		28
Other forms of sense	ory impa	airment	-	21		27		11
Titubation -	-	-	-	2		4		16
Labyrinthine sympto	oms	-	-	7		17	•••	

*64 cases seen in the survey less three "early" cases.

disseminated sclerosis. The onset is before the age of 20 and similar in the siblings, although this is not so when the inheritance is due to a dominant gene (Bell, 1939). The progressive history of spastic or ataxic weakness of the lower limbs can also occur in disseminated sclerosis. Carter, et al. (1950), investigated the clinical records of 46 cases of disseminated sclerosis in whom the diagnosis had been confirmed by autopsy. Remissions occurred in 59 per cent. of their series and weakness of the legs remitted in only 26 per cent. Deformities such as scoliosis and pes cavus may not be present in all cases of hereditary ataxia-Carmichael and Bell found no scoliosis in 4 and no pes cavus deformity in 7 out of a total of 40 cases of spastic ataxia in their English material; also 6 of the 40 showed severe mental deterioration; 7 had generally pale discs and 4 temporal pallor, without visual impairment; urinary symptoms occurred in 8 (Bell, 1939). Ophthalmoplegia of varying degrees is a well-known phenomenon in cases of hereditary ataxia. Retrobulbar neuritis, which is usually regarded as strong evidence in favour of disseminated sclerosis, has been reported in 2 members of a family suffering from hereditary spastic paraplegia. Both cases were under the age of 10 and had no other neurological symptoms or signs (Bickerstaff, 1950). Leeuwen and Van Bogaert (1949) found that in certain cases of hereditary ataxia optic atrophy shows the typical picture of "retrobulbar neuritis"; the onset of blindness may be acute and a partial remission may occur, as in Leber's optic atrophy. The condition, however, does not altogether mimic the retrobulbar neuritis of disseminated sclerosis in which the lesion is usually unilateral and remits in a matter of weeks or months, although this did happen in one of Bickerstaff's patients. Nor can the presence of a Lange curve associated with a negative Wassermann in the C.S.F. always be considered as confirmatory evidence in favour of disseminated sclerosis (Aring, 1938). Thus it would be possible to construct from the above facts a case of hereditary ataxia which would be indistinguishable clinically from disseminated sclerosis.

At this juncture it is pertinent to mention two interesting instances in the literature where pathologically there were present both the lesions of Friedreich's ataxia and disseminated sclerosis. In 1922 Mondini published a report of a young woman whose clinical history was typical of Friedreich's ataxia; the autopsy was, unfortunately, limited to examination of the cerebellum and medulla; sections showed symmetrical degeneration of the direct spino-cerebellar tracts and other tracts, also many sclerotic plaques. He was of the opinion that this was a case of Friedreich's ataxia complicated by disseminated sclerosis. More convincing is Brouwer's report in 1933 of two sisters suffering from Friedreich's ataxia. Postmortem findings in one sister showed, in addition to the findings of Friedreich's ataxia, lesions resembling the plaques of disseminated sclerosis.

There were 18 cases of hereditary spastic ataxia among those notified, and in Table 3 we have compared them on the basis of symptomatology with the familial and probable groups. The number of cases in each of the three groups is so dissimilar that it is not possible to draw any significant conclusions, but the table does illustrate some of the points of differential diagnosis between disseminated sclerosis and the hereditary ataxias mentioned above; also it shows that the distribution of symptoms in the familial group is similar to that in the probable group. It has not always been possible, however, to make a firm diagnosis, and below are two examples which were excluded from the series :—

Female, Mrs. G., born 1902: 1914, at the age of 13, numbness and weakness of right hand for a few months; 1918, diplopia on two occasions, each lasting one week; 1920, the vision of the right eye was hazy for two weeks; 1928, dizziness and numbness of the legs and back which cleared up rapidly; 1939, gradually increasing weakness of the right leg and staggering. Since 1944 left leg also affected. Bedridden since 1946. The right foot had been deformed from birth; 1949, examination showed marked right-sided clubfoot deformity; visual acuity R=J.12, L=J.2; pallor of both discs; weakness and hypotonia of arms, especially the right; wasting of the small muscles of the right hand; slight intention tremor left finger-nose test, limited voluntary movements left leg only; legs contracted, left extended, right semi-flexed; extensor plantar responses; all deep tendon reflexes were unobtainable; abdominal reflexes absent; vibration sensation absent throughout; muscle joint sensation absent in toes.

Family History : Mother alive and well, aged 74. Father died of "ulcer" at 35. Mother's cousin in U.S.A. has a neurological condition of 20 years' duration. Sister, Mrs. M. D., as below.

Female, Mrs. M. D., born 1896: 1928, aged 32, blindness in right eye for a few months which recurred in 1935; 1945, increasing weakness left leg; 1948, admitted to Royal South Hants Hospital, then complained of precipitancy of micturition. On examination—Slight nystagmus to right and left; bitemporal pallor of discs; weakness of both legs, especially the left; generalized hyperreflexia; absent abdominal reflexes; extensor plantar responses; absent vibration sensation in legs; postural sense diminished left leg.

C.S.F.: Cells 2; protein 50 mgms. per cent.; globulin : faint trace; W.R. : negative; Lange 2344432100.

Later bedridden and paraplegia in flexion developed and she died in April, 1952.

To summarize : Two sisters with history in keeping with disseminated sclerosis. One had a congenital clubfoot deformity and the tendon reflexes were unobtainable, although contractures could account for the absent knee and ankle jerks. For these reasons this family was excluded from the series.



(iii) 1.—*Case J. A., male, born 1911*: 1934, "useless" right hand for three weeks; 1944, blurred vision for three weeks; 1949, gradually increasing stiffness of legs and hesitancy of micturition with partial remission after two years. Examination 1950: Cranial nerves normal; right hand slightly clumsy; spastic weakness of legs with generalized hyper-reflexia; abdominal reflexes absent on left side; plantar reflexes extensor; vibration sensation absent in legs; position sensation slightly diminished in toes. C.S.F.: White cells 3; protein 55 mgms. per cent.; globulin trace; W.R.: negative; Lange 4432100000.

(ii) 1.—*Case J. C., male, born 1880*: 1936, in the National Hospital, Queen Square, London, under the care of Dr. Gordon Holmes with a diagnosis of subacute combined degeneration of the spinal cord. Two years' history of aching and stiffness of the legs. Examination at that time showed no abnormality in the cranial nerves; slight inco-ordination in the finger-nose test of both arms; there was spastic weakness of both legs with sustained knee clonus, absent ankle jerks and extensor plantar reflexes; vibration sensation was diminished in the legs. R.B.C. count: 4.84 million; hb. 90 per cent.; colour index 0.93; F.T.M. showed achlorhydria. W.R.: negative. C.S.F. normal.

Patient died in 1950 in the Home and Hospital for Jewish Incurables, where he was considered to be suffering from disseminated sclerosis.

(ii) 2.—*Female, born 1882*: According to her doctor, she has been suffering from "spastic paralysis" of 15 years' duration, but is able to walk with sticks. No other signs. Mentally normal. No nystagmus. No intention tremor. No dysarthria.

(ii) 3.—Female, born 1890 : A progressive illness since the age of 40; now bedridden.

(i) 1 and 2.—Grandparents : Polish and first cousins.

Although case J. A. could be diagnosed as disseminated sclerosis, his uncle, J. C., in the early stages of his condition, was diagnosed as subacute combined degeneration of the cord, and it is most unlikely that the condition was disseminated sclerosis in view of the absent ankle jerks. For this reason, this family was discarded from the series.

OTHER FAMILIAL NEUROLOGICAL DISEASES.

There are other less common familial neurological disorders which can simulate disseminated sclerosis. Ferguson and Critchley (1929) described a form of hereditary ataxia resembling disseminated sclerosis, the unusual features being limitation of upward gaze, exophthalmos and parkinsonism. In 1907 Holmes described a form of familial degeneration of the cerebellum. In another paper, he reviewed and classified cerebellar disease. He did not consider olivo-pontocerebellar atrophy to be hereditary or familial. However, in a more recent paper Critchley and Greenfield (1948) found a few familial cases of olivo-ponto-cerebellar atrophy in the literature, and distinguished this condition from cerebello-olivary degeneration, which was mainly familial, on pathological grounds.

Ferraro (1927) reported a familial form of encephalitis periaxialis diffusa occurring in two brothers and one sister, who were clinically diagnosed as disseminated sclerosis. In each case the onset of the disease occurred in the third

 TABLE 4.

 All cases in this table are counted as affected whether "probable,"

 "possible," or "early." The table includes all sibships

 whether no, or one, parent is affected.

Family	No. of		Males			Females		Total		
Size	Families	A	U	Т	A	U	T	A	U	T
1	36	19		19	17		17	36		36
2	60	28	29	57	34	29	63	62	58	120
3	70	31	66	97	39	74	113	70	140	210
4	100	47	149	196	56	148	204	103	297	400
5	86	48	181	229	45	156	201	93	337	430
6	97	41	262	303	63	216	279	104	478	582
7.	75	40	229	269	40	216	256	80	445 ,	525
8	64	21	238	259	45	208 ,	253	66	446	512
9	42	17	154	171	31	176	207	48	330	378
10	21	8	98	106	14	90	104	22	188	210
11	6	3	32	35	4	27	31	7	59,	66
12	6	2	31	33	4	35,	39	6	66	72
13	4	1	27 ,	28	3	21	24	4	48	52
14	1	1	4	5		9	9	1	13	14
Total	668	307	1500	1807	395	1405	1800	702	2905	3607

 $A = Affected. \qquad U = Unaffected. \qquad T = Total.$

decade. He discussed the possible relationship of this condition to Pelizæus-Merzbacher disease.

THE PROBLEM.

When, in disseminated sclerosis, as in many other conditions, the familial incidence of the condition is definite but low, many problems arise in interpretation. It is clear that the proportion affected in the sibships of the propositus is too low to be interpreted as the expression of a single gene. By the word 'propositus' we mean the index case or case which brought the family to our attention. The minimum proportion to be expected on a single gene hypothesis would be 1 in 4, whereas the actual expression is about 1 in 50 (allowing for the fact that the method of ascertainment was by one affected sib in the sibship). To satisfy any single gene hypothesis it would therefore be necessary to postulate that the gene was only expressed in less than 10 per cent. of the people who had the gene. The

TABLE 5. All cases in this table are affected whether they are "probable," "possible," or "early." The table excludes sibships where a parent is affected.

Family	No. of		Males			Females	5	Total		
Size	Families	A	U	Т	A	U	T	A	U	T
1	36	19		19	17		17	36		36
2	59	27	28	55	34	29	63	61	57	118
3	69	30	64	94	39	74	113	69	138	207
4	96	43	145	188	56	140	196	99	285	384
5	85	47	180	227	44	154	198	91	334	425
6	96	41	257	298	62	216	278	103	473	576
7	74	39	228	267	40	211	251	79	439	518
8	63	20	235	255	45	204	249	65	439	504
9	41	17	151	168	29	172	201	46	323	369
10	21	8	98	106	14	90	104	22	188	210
11	6	3	32	35	4	27	31	7	59	66
12	6	2	31	33	4	35	39	6	66	72
13	4	1	27	28	3	21	24	4	48	52
14	1	1	4	5		9	9	1	13	14
Total	657	298	1480	1778	391	1382	1773	689	2862	3551

more usual methods of expressing this would be to say that the gene had less than 10 per cent. penetration or that less than 10 per cent. of the susceptible genotypes showed the trait.

It is always tempting to proceed logically at this stage and to say that perhaps two genes are involved, and by juggling with possible combinations of one or two dominant and/or recessive genes to demonstrate that one's own observations fit in with some theoretical hypothesis. For many reasons this is a dangerous exercise. However, proceeding along somewhat different lines from our original observations of undue concentration of cases within sibships, it is more reasonable to argue as follows :—

First, we are going to consider the sibships in which the cases occur, and we want to know whether there is undue concentration of cases in these sibships. There will always be at least one case in each sibship, because each was 'ascertained' by a propositus or index case. Therefore we must allow for this or we should never in any condition arrive at an incidence in sibships so ascertained of less than one divided by the mean sibship size. Or to reduce to absurdity in another way—if all the families consisted of two sibs we should have a minimum of 50 per cent. affected. To be accurate, as, for example, if we were trying to fit our observations to a simple ratio, we should have to make a separate adjustment for each family

ΤA	۱B	LE	6.

All cases in this table are regarded as affected as in Tables 4 and 5.

Family	No. of		Males			Females			Total	
Size	Families	A	U	Т	A	U	Т	A	U	Т
1		_			_					_
2	1	1	1	2	—	— .		1.	1	2
3	1	1	2	3		i —		1	2	3
4	4	4	4	8	_	8	8	4	12	16
5	1	1	1	2	1	2	3	2	3	5
6	1		5	5	1		1	1	5	6
7	1	1	1	2		5	5	1	6	7
8	1	1	3	4		4	4	1	7	8
9	1		3	3	2	4	6	2	7	9
Total	11	9	20	29	4	23	27	13	43	56

This table includes only sibships where one parent is affected.

size. This is hardly worth while. Nevertheless, we have set out the data in tables by family size (a) in order to make it clear that we recognize the point; and (b) so that it may be possible, if desired, to make further calculations (Tables 4-9).

We do not know the frequency of the genotype which is often, but not constantly, expressed as disseminated sclerosis. However, if we presume that the disease is the expression, precipitated by environmental factors of a specific genotype, then we should expect that the same genotype would occur more frequently in the sibs of affected persons than in the general population. If we exclude from our calculations the affected individual by whom we identified the family, all the others have independently the same chance of being affected and the fraction, affected sibs divided by total sibs, will give the incidence in the sibs, which can then be compared with that in the general population.

TABLE 7.

ALL PROBABLE SIBSHIP

i.e., where at least one affected person was regarded as a "probable" case. "Possible" or "early" cases are counted as "unaffected" in this table. It includes all families, whether parent affected or not.

Family	No. of		Males			Females			Total		
Size	Families	Α	U		A	U	Т	Α	U	T	
1	26	15		15	11		11	26		26	
2	47	22	24	46	26	22	48	48	46	94	
3	43	18	40	58	25	46	71	43	86	129	
4	68	28	100	128	40	104	144	68	204	272	
5	61	34	126	160	31	114	145	65	240	305	
6,	59	19	160	179	43	132	175	62	292	354	
7	46	22	143	165	25	132	157	47	275	322	
8	46	14	170	184	33	151	184	47	321	368	
9	33	13	127	140	22	135	157	35	264	297	
10	12	4	57	61	9	50	59	13	107	120	
11	3	1	16	17	2	14	16	3	30	33	
12	4	1	18	19	3	26	29	4	44	48	
13	4	1	27	28	3	21	24	4	48	52	
14	1	1	4	5 		9	9	1	13	14	
Total	453	193	1012	1205	273	956	1229	466	1968	2434	

TABLE 8.

PROBABLE SIBSHIP—excluding those where a parent was affected.

As in Table 7, "possible" or "early" cases are counted as "unaffected" in this table

unamected in this table
unaffected in this table

Family	No. of		Males			Females	6	Total		
Size	Families	A	U	Т	A	U	T	A	U	Т
1	26	15		15	11		11	26		26
2	46	21	23	44	26	22	48	47	45	92
3	43	18	40	58	25	46	71	43	86	129
4	66	26	96	122	40	102	142	66	198	264
5	61	34	126	160	31	114	145	65	240	305
6	58	19	155	174	42	132	174	61	287	348
7	46	22	143	165	25	132	157	47	275	322
8	46	14	170	184	33	151	184	47	321	368
9 :	33	13	127	140	22	135	157	35	262	297
10	12	4	57	61	9	50	59	13	107	120
11	3	1	16	17	2	14	16	3	30	33
12	4	1	18	19	3	26	29	4	44	48
13	4	1	27	28	3	21	24	4	48	52
14	1	1	4	5		9	9	1	13	14
Total	449	190	1002	1192	272	954	1226	462	1956	2418

THE DATA USED IN THE GENETIC ANALYSIS.

The Sibships of the Propositi.

The genetic analysis considers in all 668 sibships with 702 cases in the sibship of the propositi.

The reasons why the genetic analysis considers 702 affected sibs in the sibship of the propositi while the epidemiological analysis considers 700 cases are as follows :----

1. In the epidemiological analysis are included 20 cases excluded from the genetic analysis for the following reasons :

(i)	Sibship size not known	No. of Cases 13
(ii)	Cases not in the sibship of the propositus, although they were relations of the propositus	7

2. In the genetic analysis are included cases where sibs were regarded as affected, but they were either dead or they were not seen by the observers. These totalled 22 cases, i.e., there are in all 2 more (22 minus 20) cases considered in the genetic analysis. Siblings under 15 years, or who did not survive 15 years, are omitted.

TABLE 9.

PROBABLE SIBSHIP—only those with one parent affected are included in this table. As in Table 7, "possible" or "early" cases are counted as "unaffected" in this table.

Family	No. of	Males				Females		Total		
Size	Families	A	U	Т	A	U	Т	A	U	Т
1			_							
2	1	1	1	2		—	—	1	1	2
3										
4	2	2	4	6		2	2	2	6	8
5								_		
6	1		5	5	1		1	1	5	6
									·	
Total	4	3	10	13	1	2	3	4	12	16

TABLE 10.

SUMMARY.

		Neit	her Pare	NT AFFE	CTED		
Clinical	No. of		Males			Females	
cation	Sibships	A	U	T	A	U	Т
"Probable" Sibships	449	190	1002	1192	272	954	1226
All Sibships	657	298	1480	1778	391	1382	1773

		On	e Parent	AFFECT	ED				
Clinical Classifi-	No. of		Males			Females			
cation	Sibships	А	U	T A U 0 13 1 2	Т				
"Probable" Sibships	4	3	10	13	1	2	3		
All Sibships	11	9	20	29	4	23	27		

					Total					
Clinical Classifi-	No. of		Males			Females			Total	
cation	Sibships	A	U	Т	A	U	Т	Α	U ·	Т
"Probable" Sibships	453	193	1012	1205	273	956	1229	466	1968	2434
All Sibships	668	307	1500	1807	395	1405	1800	702	2905	3607

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Probable sibships are those where at least one case which is classed as probable occurs. In this line any "possible" or "early" cases which occur in sibship or parent are counted as unaffected.

All sibships includes every case occurring in the sibship of the propositus : i.e., "probable," "possible," and "early" cases are all counted as affected. Similarly, an affected parent, by that definition, puts the appropriate sibship in the "One parent affected" column.

11.	
TABLE	

PARENTAL CONSANGUINITY.

	Presen	T SERIES	MA	THERS	CURTIC	JS (1933)	Bell	(1940)	Prati	r (1951)
	No.	Per Cent.	No.	Per Cent.	No.	Per Cent.	No.	Per Cent.	No.	Per Cent.
Number of Families	*558	100.00	670	100.00	106	100.00	632	100.00	134	100.00
No Consanguinity	538	96.42	668	99.70	100	94.34	624	98.73	131	97.76
First Cousin Marriages	9	1.08	I		3	2.83	9	0.95	5	1.49
Second Cousin Marriages	9	1.08	7	0.30						
Paternal Grandparents First Cousins	4	0.72	1		ŝ	2.83	7	0.32		0.75
Other Relationships	4	0.72	1				-			
Total Incidence of Consanguinity -	20	3.58	5	0.30	و	5.66	8	1.27	3	2.24

*Of the 668 sibships considered, information was available only in respect of 558 sibships.

INCIDENCE IN THE SIBS OF THE PROPOSITI.

In the 668 sibships there were 3,607 sibs, and of these 702 were affected. Ignoring the propositi, there were 34 affected sibs in 2,939 sibs of the propositi at risk, i.e., 1.15 per cent. (Table 4). Where neither parent was affected there were 657 sibships, having in all 3,551 sibs, and of these 689 were affected. Thus, 32 in 2,894 sibs at risk were affected when the propositi are ignored, or 1.11 per cent. (Table 5). Where one parent was affected, in 11 sibships there were 56 sibs, and in all 13 sibs were affected, so that 2 of 45 sibs of the propositi or 4.44 per cent. were affected (Table 6).

	5	Sex I	Distrib Sib F	ution i Pairs ai	in Affe nd Thre	ected S	Sibs.		
Sib S Grou	e x ps		No. of Families	5	М		F		Total
ΜΜ	-	-	—						
ΜF	-	-	13		13		13		26
FΓ	-		9				18		18
2M F	-	-	2		4		2		6
M 2F	-	-	1		1		2	•••	3
3M	-	-	1	•••	3				3
3F	-	-							
	Total	-	26		21		35		56

TABLE 12.

Similar calculations are made for those sibships where there was at least one probable case (Tables 7-9). There were 13 in 1,981 sibs, or 0.66 per cent., affected. Where neither parent was affected, 13 in 1,969 sibs, or 0.66 per cent., and where one parent was affected there were no sibs affected in 12, or 0 per cent., but in this instance the small number of sibships make it impossible to draw any conclusions. It should be stressed that, in the probable sibships, all possible and early cases were counted as unaffected in these calculations. Tables 4-9 set out the data for family size and the figures are summarized in Table 10. It will be noted that the number of very large families, 10 or over, is small, but there appears to be a trend for the number of multiple cases within sibships to increase between family sizes 2 and 9 where the number of sibships is considerable.

The incidence in the sibs of the propositi, 1.15 per cent., is greater than the prevalence rate in the general population in Northern Ireland when all age and diagnostic groups are included, 0.072 per cent. It is also greater than the highest rate of 0.182 per cent,—that of County Fermanagh in the age group 40-59 years. (Allison and Millar, 1954.) When we confine our attention to the incidence in the probable group, or 0.66 per cent., it is again greater than the prevalence rate of cases in the general population, 0.05 per cent., even 0.124 per cent., the highest rate again in County Fermanagh in the age group 40-59 years. It would therefore be fair to say that the incidence in the sibs of the propositi is somewhere between 5 and 15 times greater than the prevalence rate in the general population. Although

TABLE 13.

Sex linkage information from Consanguinity Data. Sex of parent of offspring of first-cousin marriages with sex of paternal grandparent.

			Rel	ationship through Father's
Serial No.		Sex of Patient		Father or Mother
164		F		Mother
251	•••	F		Mother
267		Μ		Mother
597		\mathbf{M}	•••	Father
708		F	•••	Mother
830		F		Mother

these figures are impressive, it is difficult to be sure that they are technically significant, a fact not always taken into account. There are two main difficulties in making a numerical comparison. The first is the size of the sampling error involved, and secondly, the difficulty in making standardizations for age.

CONSANGUINITY.

In 558 of the 668 sibships considered, information about consanguinity of the parents of the propositus was collected (Table 11). The parents of the propositus were full cousins in six instances and second cousins in a further six instances. There was believed to be a common ancestor of parents several generations back, the relationship not being quite clear in four instances. That is, in 16 sibships there was some degree of consanguinity of parents. In addition, in four instances

the parents of the propositus' father were full cousins. Full-cousin consanguinity of parents, therefore, occurred in 6 of 668 sibships, or 0.90 per cent.; if the sibships where no information was available are excluded, 6 of 558 sibships, or 1.08 per cent.

No reasonable control figures are available for comparison as Bell's (1941) figures for England and Wales could not safely be used. Mathers (1952) found that of 670 married adults questioned in casualty department of the Royal Victoria Hospital, Belfast, two were married to a second cousin. It cannot be said that consanguinity is unduly common in the parents of disseminated sclerotic patients, especially when compared with, for instance, the figure 21.7 per cent. found in the parents of patients suffering from the myopathies (Stevenson, 1953). Table 11 also sets out the consanguinity rates in disseminated sclerosis in other papers. There is reasonable agreement.

SEX INCIDENCE AND ASSOCIATIONS.

The sib pairs and threes in the 26 sibships are shown in Table 12. These figures do not suggest any tendency for one sex to be more affected than the other. The information about the sex of the patient, relevant parent and grandparent, where there was consanguinity, is shown in Table 13. Here the numbers are small, and speculation about partial sex linkage seems too hazardous to warrant discussion, although our findings are similar to those shown in Pratt's Table 6 (Pratt, 1951).

DISCUSSION.

This is the first time that the familial and general population rates of disseminated sclerosis have been ascertained in the same area at the same time. We have also been fortunate in that there has been a census of the population in Northern Ireland in 1951 during our investigation. We feel confident in stating that there is a familial incidence in this disease and can answer Mackay's question in the affirmative— "However, the precise question we wish answered is whether the incidence of familial multiple sclerosis is greater than the incidence of the disease in the general population" (Mackay, 1950). The figures, although conclusive, are small, and the familial factor cannot be the only one; like many other diseases, there are both genetic and environmental factors.

There are no known generally accepted environmental factors, but recently there has been agreement that a familial factor exists (Mackay, 1950; Pratt, et al., 1951). This is at present the only widely accepted factor in the ætiology of the disease. In the Middlesex Hospital series the incidence in the sibs was 0.82 per cent.; in the Curtius and Speer series (1937) 0.9 per cent.; these percentages correspond very closely to the figure in this series which lies between 0.65 per cent. and 1.15 per cent.

In the past a common environment or exposure to the same "toxin" has been used to explain the occurrence of the familial cases; however, if this were true, one would expect a higher incidence in husbands and wives than in the general population. There was no instance of conjugal disseminated sclerosis in this series, and we have found only two instances in the literature (Steiner, 1938). Table 14 shows the environmental factors in our families, where these were known, and in four instances there were no common environmental factors.

No paper on the familial aspects of disseminated sclerosis would be complete without mentioning the literature on twins suffering from this disease. Strangely, we had only one instance of an affected dizygotic twin. Table 15 sets out the number of monozygotic twins affected in the literature. Dizygotic twins are not included, as the genetic risk is no greater than that of siblings. From the table it will be seen that 30.8 per cent. of the sibs at risk were affected. This is strong supporting evidence that there is a genetic factor, and, again, that this cannot

TABLE 14.

Environmental Factors in the A	FFECTED MEMBERS	6 OF 32	FAMILIES.
The period of common environme one member	nt ceased after the	e onset ir	n - 6
The period of common environme in both members	nt continued after	the onse	t - 10
The period of common environme in all three members	nt continued after	the onse	t - 2
The period of common environmen all affected members	t ceased prior to th 	e onset in	- 10
No common environmental factors Serial Numbers—278, 414, 489,	s— and 339 -	-	- 4

be the only factor. It should be stated, however, that, by modern standards, the evidence that the twins were monozygotic was in some instances lacking.

There are other conditions where genetic factors play a part, of much the same order as in disseminated scleroris. In juvenile rheumatism 5.03 per cent. of the sibs at risk are affected (Stevenson and Cheeseman, 1953). Harris (1951) found 4.3 per cent. of the sibs affected in diabetes mellitus. Also in diabetes mellitus Steinberg, et al. (1952), found 4.7 per cent. incidence in the sibs of the propositi where neither parent was affected; 11.4 per cent. where one parent was affected, and 16 per cent. where both were affected. In psoriasis, the figures were 2.45 per cent. where neither parent was affected and 9.0 per cent. where one parent was affected (Steinberg, et al. (1951). Stamos (1940), in a series of 645 cases of pernicious anæmia, found the familial incidence to be 7.9 per cent.

TA	۱B	LE	15

Both One Author affected unaffected Remarks (a) Onset 19, died 28 (male). Legras (1934) 1 _ . . . (b) Onset 24, died 27 (male). Kranz (1936) 1 Microcephalic observed until 22 years . . . (male). 1 Jentsch (1937) . . . Onsets 19 and 31 (male). 1 Onsets 24 and 48 (male). Curtius & Speer -. . . (1937)Voss (1937) -1 Observed until 42. . . . Isenschmid & Olloz 1 Onsets 24 and 26 (male). . . . (1939) $\mathbf{2}$ Schaltenbrand (1943) ----. . . Williams (1946) -1 ____ Onset at 30; no details of second twin . . . (female). 1 Jequier (1949) Onsets 30 and 36 (female). . . . 1 Male. Reese (1950) . . . _ 2 (a) Onset 25 (female); observed for Pratt (1951) _ . . . two years. (b) Onset 47 (male); observed two years. 12Thums (1951) 1 . . . 18 8 TOTAL -. . .

CASES OF DISSEMINATED SCLEROSIS IN MONOZYGOTIC TWINS.

This table is a modification of Pratt's Table 3 (Pratt, 1951).

SUMMARY.

In Northern Ireland we have found 44 families with two or more members affected out of a total of 668 families, an incidence of 6.58 per cent.

The incidence of the disease in the sibs of the propositi is 5 to 15 times greater than the prevalence rate in the general population.

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APPENDIX X.

Case histories of families 1-23 inclusive, in which we have examined two or more affected members.

Case histories of families 24-29 inclusive, in which we examined one member, and the evidence from another source was sufficiently strong to warrant a firm diagnosis in the other.



1.—Serial No. 644, female, born 1908: 1942, aged 34, gradually increasing weakness right leg, intermittent attacks of diplopia; 1949, increasing difficulty in starting micturition and weakness of arms. Examination 1951, bilateral pallor of discs, visual acuity normal, spastic weakness of legs, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses, impaired vibration and muscle joint sensation in legs. (Probable D.S.)

2.—Serial No. 414, female, born 1920: 1946, aged 26, headache, vomiting and generalised paræsthesiæ; examination, small central scotoma right eye, nystagmus to right, ataxic weakness of right arm and legs, astereognosis right hand, impairment of sensation to pain and light touch over trunk and legs, generalised hyperreflexia, absent abdominal reflexes; complete recovery in two months; 1949, sudden paralysis of arms and legs, with retention of urine; complete recovery in one month. Examination, bilateral temporal pallor of discs, good visual acuity, total quadriplegia except slight movements of fingers, flexors of knees and dorsiflexors of feet, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses; 1952, paræsthesiæ in limbs for two weeks, power normal and plantar reflexes flexor. (Probable D.S.)

Cousins, no common environmental factor.



1.—Serial No. 551, female, born 1904: 1951, aged 46, weakness left leg, tendency to drop things out of left hand with slight weakness of left arm, curious paræsthesiæ in head and shoulders; no previous episodes. On examination, slight weakness left arm and leg, rapidly repeated movements not quite normal left hand, hyperreflexia and doubtful extensor response on left side. (Early D.S.)

2.—Serial No. 78, female, born 1913: 1939, aged 26, "useless" right arm, difficulty in appreciating the nature of objects in the hand when the eyes were closed; this symptom lasted four months; 1940, weakness of right leg, which gradually got worse, especially after the birth of her second child with slight precipitancy of micturition. Examination 1948, temporal pallor left disc, V.A. 6/6, 6/6, spastic weakness right leg, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, vibration diminished right leg; 1950, weakness of legs improved and able to carry on with household duties. (Probable D.S.)

Two sisters, same environment.



1.—Serial No. 625, male, born 1921: 1940, aged 19, dragging right leg, progressive and spreading to other leg, temporary cold feeling in legs and slight weakness of hands; 1951, examination, pallor of both discs, V.A.R. J2, V.A.L. J12, very slight intention tremor both arms, legs ataxic and slight weakness right leg, generalised hyperreflexia, extensor plantar responses, vibration sensation absent and muscle joint sensation defective in legs. (Probable D.S.)

2.—Serial No. 313, female, born 1922: 1943, aged 21, pain and loss of vision left eye, with partial recovery in four months; 1944, diplopia, weakness and paræsthesiæ of legs, with partial recovery; 1950, increasing weakness and ataxia legs. Examination 1951, euphoric, bilateral optic atrophy, with impairment of visual acuity, weakness of left external rectus, horizontal nystagmus on right lateral gaze, intention tremor of arms, weak spastic ataxic legs, generalised hyper-reflexia, absent abdominals, extensor plantar responses, impaired vibration and muscle joint sensation in legs. (Probable D.S.)

3.—Case M. McC., male, born 1925, died 1947, diagnosed as disseminated sclerosis.

All three siblings lived at same address.



1.—Case J. A. H., male, born 1913: 1944, aged 31, numbness and awkwardness of the right leg for one month, with complete recovery; 1951, rapid onset of numbness and weakness of right leg progressing in a matter of days to ataxia so that he was unable to walk. Examination then showed weak ataxic legs, with extensor plantar reflexes and absent abdominal reflexes. Subjective complete recovery in one month. Examination then showed considerable improvement, the plantar reflexes being doubtfully extensor, vibration sensation absent and muscle joint sensation impaired in the legs. (Probable D.S.)

2.—Serial No. 18, male, born 1915: History not very reliable, but certain facts are available. The illness began 1934-38. He was a patient under Professor Sir W. W. D. Thomson in the Royal Victoria Hospital in 1940, when he complained of girdle pains around his abdomen, vertigo, numbness of the hands and ataxic gait. There was also a note that he had had "poliomyelitis" at the age of 18 months, with a residual weakness of the left leg. Examination 1949, bedridden, dysarthric, nystagmus to right and left, right-sided facial weakness of upper motor neurone type, tongue protruded to right, marked ataxia of upper limbs, with intention tremor, legs paraplegic in flexion, incontinent of urine, no definite sensory changes. Died 1950. (Probable D.S.)

3.—Serial No. 19, male, born 1919: 1942, aged 23, frequency and precipitancy of micturition, no improvement; 1943, gradually increasing failure of vision of both eyes until unable to read small print; 1945, increasing weakness and ataxia of legs. Examination 1950, bilateral optic atrophy, with central scotomata. V.A.R. 6/60. V.A.L. 6/90. Monocular coarse horizontal nystagmus on right lateral gaze, vertical nystagmus on upward gaze and slight rotatory nystagmus in the central neutral position, slight ataxia in arms, ataxia in legs, all modalities of sensation impaired slightly in legs, generalised hyperreflexia, absent abdominal reflexes and extensor plantar responses. (Probable D.S.)

Three brothers-same environment until 1935.



1.—Serial No. 912, male, born 1925: 1945, aged 20, gradual onset of frequency and urgency of micturition, no remission; 1946, diplopia for two months; 1951, "useless" right arm and weak right leg for two months, with partial recovery; 1952, gradually increasing weakness left arm and leg. Examination 1952, bitemporal pallor. V.A.R. and L J2, coarse horizontal nystagmus on right lateral gaze, slight intention tremor of arms, dysdiadokokinesia left arm, slight ataxic weakness left leg, left plantar reflex extensor, brisk tendon reflexes, absent abdominal reflexes, impaired vibration sensation in legs. (Probable D.S.)

2.—Case J. McN., male, born 1897: 1940, aged 43, gradually increasing weakness left leg, paræsthesiæ left hand; variable course, but no clear-cut remission; 1948, weakness spread to right leg; 1950, slowness in micturition. Examination 1952, slight pallor left disc, horizontal monocular nystagmus on left lateral gaze, weakness of legs, increased reflexes in legs, absent abdominal reflexes, extensor plantar reflexes. (Possible D.S.)

Father and son, same environment.

The following case, not included in the series, has recently been seen, and is the eldest child of J. McN.'s only sister.

Case Mrs. R., female, born 1930: 1950, loss of sight right eye for three weeks, with complete recovery. March, 1953, paræsthesiæ in all limbs, vertigo, transient loss of sight right eye, diplopia and weakness of limbs. On examination, nystagmus, generalised weakness, increased tendon reflexes, extensor plantar reflexes, absent abdominal reflexes, impaired postural and vibration sensation in arms. (Probable D.S.)



1.—Case, Mrs. McD., female, born 1882, died 1941: 1926, gradually increasing weakness of legs until, by 1930, unable to walk. (Possible D.S.)

2.—Serial No. 489, male, born 1895 : 1943, aged 48, difficulty in concentrating and lapses of memory; 1946, difficulty walking and retention of urine. Examination 1947, dementia, small contracted unequal pupils, with sluggish reaction to light and convergence, weakness right arm, spastic ataxic legs, with hyperreflexia, absent right abdominal reflexes, extensor plantar responses; 1948, made remarkable spontaneous remission; 1949, Sir Russell Brain diagnosed disseminated sclerosis. April, 1949, slight pain, with loss of vision of right eye, with recovery in one month. On examination, pallor of both discs, jaw jerk increased, intention tremor and slowing of rapidly alternating movements of arms. Abdominals absent. Unsustained patellar and ankle clonus, generalised hyperreflexia, plantar reflexes extensor, loss of sense of passive movement right great toe. (Probable D.S.)

3.—Serial No. 65, male, born 1904: 1924, aged 20, pain right buttock and leg diagnosed as "sciatica" for two years; 1932, sensation like "belt" and paræsthesiæ in legs, two months' duration; 1936, dragged left leg for two months, with partial recovery; 1937, diplopia for one month and since intermittently; 1939, vomiting, followed by gradual weakness both legs lasting two months, with partial recovery; 1946, loss of power of legs for one month, with partial recovery; 1947, feeling "as if walking on wooden legs" for one month, with partial recovery; 1949, increasing weakness of legs. Examination 1949, walking quite well with stick, monocular nystagmus to right and left, temporal pallor left disc, visual acuity normal, slight weakness and spasticity of legs, absent abdominal reflexes, generalised hyperreflexia and extensor plantar responses, impaired muscle joint and vibration sensation in legs. (Probable D.S.)

Nephew and uncle, different environments.



1.—Serial No. 229, female, born 1907 (history not very reliable as memory impaired): 1947, aged 40, "nervous breakdown," weeping, depressed, ataxia, and precipitancy of micturition; complete recovery in four months except for persistence of precipitancy of micturition; 1949, gradual weakness left leg, ataxic gait and clumsy left arm. Examination 1950, dementia, euphoria, monocular nystagmus on right and left lateral gaze, weak handgrips, slight ataxic left arm, two-point discrimination impaired left hand, ataxic weakness of legs, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, impaired vibration sensation in legs. (Probable D.S.)

2.—Serial No. 830, female, born 1910: History vague and indefinite, but has dragged left leg "for years." Examination 1951, bitemporal pallor of discs, generalised hyperreflexia, left knee and ankle jerks brisker than right, left plantar reflex extensor. (Possible D.S.)

Two sisters, same address until 1932.



1.--Serial No. 484, female, born 1926: 1947, aged 21, paræsthesiæ right foot, and twice had fallen because of tripping with the right foot; 1948, "heavy sensation" left arm; all the symptoms cleared up in 1949. Examination 1950, no abnormal neurological signs. (Early D.S.)

2.—Serial No. 485, male, born 1929: 1944, aged 15, weakness and paræsthesiæ of legs for three months; 1949, gradual onset weakness right leg, with complete recovery in four months. Examination 1950, no abnormal neurological signs except that both knee jerks increased, especially the right. (Early D.S.)

Brother and sister, same address.



1.—Serial No. 773, male, born 1898: 1935, aged 37, blurred vision for one month; 1939, trailing right leg after a long walk. Examination 1951, pallor both discs, V.A.L. J4, V.A.R. J4, horizontal and slightly rotatory nystagmus to left, bilateral ptosis, slight weakness of arms with slight ataxia, paraplegia in flexion, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, vibration and position sensation, impaired in legs. (Probable D.S.)

2.—Serial No. 924, female, born 1903: 1942, aged 39, numbness left side of face and tongue for one year; some months later, following lumbar puncture at National Hospital, Queen Square, developed weakness and pains in legs for two years, with complete recovery; 1947, dragging right leg, partial recovery after a few months; 1949, "film" over both eyes for three months. Examination 1951, creamy pallor of both discs, V.A.L. J2, V.A.R. J4, weakness right leg, with slight ataxia, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses, vibration sensation absent in legs. (Probable D.S.)

Brother and sister at same address until 1934.



1.—Case M. G., female, born 1899: No detailed history available. Examination 1949, euphoria, coarse horizontal nystagmus to right and left, generalised hyperreflexia, ataxic gait, extensor plantar reflexes, bilateral ankle clonus. (Possible D.S.)

2.—Serial No. 627, male, born 1905 : 1939, aged 34, legs became weak over a period of three weeks, retention of urine and paræsthesiæ in legs, had to retire to bed, gradual improvement, and in four months could walk again with only slight residual disability. Examination 1949, spastic ataxic paraparesis, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, light touch and vibration sensation impaired in legs (had been investigated fully in Mater Hospital, Belfast, in 1945, with negative results). (Possible D.S.)

3.—Serial No. 940, male, born 1908: 1920, aged 12, sudden loss of power both legs for two months, confined to bed with gradual and nearly complete recovery, but right leg tended to drag when he walked far; 1934, onset chronic backache. Examination 1948, no abnormal findings except for nystagmoid jerks on extreme lateral gaze. (Possible D.S.)

Two brothers and one sister, similar environment until 1920.



1.—Serial No. 274, male, born 1884: 1903, aged 19, weakness of all limbs and unsteadiness of gait for eighteen months (recovery may not have been complete); 1930, gradual onset of weakness of limbs and unsteadiness of gait, with partial recovery after three years; 1931-45, diplopia at intervals; 1931, sudden urgency of defæcation. Examination 1950, bilateral optic atrophy, visual acuity two-inch print, nystagmoid jerks to the right, increased knee jerks, doubtful plantar reflexes, vibration sensation absent left leg. (Probable D.S.)

2.—Serial No. 273, male, born 1919: 1949, aged 30, gradual progressive onset of stiffness in legs and ataxia, followed in two months by intermittent diplopia; 1950, urgency of micturition. Examination 1950, temporal pallor left disc, slight ataxia of arms, weakness and spastic ataxia of legs, increased tendon reflexes, absent abdominal reflexes, extensor plantar responses, knee and ankle clonus. (Possible D.S.)

Father and son, same environment until 1938.

(12)



1.—Serial No. 58, male, born 1891: 1923, aged 32, weakness left leg, no remission; 1925, sudden dimness of vision left eye for one month; 1933, gradual weakess left arm. Examination 1951, pallor left disc, with impaired vision, weak spastic left arm, right arm slightly weak, spastic paraplegia in extension, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses. (Probable D.S.)

2.—Serial No. 51, female, born 1897: 1937, aged 40, pains and weakness of legs suddenly, following the birth of second son; the pains disappeared in two months, but the legs gradually became weaker; 1947, bedridden, urgency of micturition and onset of mental deterioration. Examination 1949, pallor of both discs, V.A. 6/9, 6/9, horizontal nystagmus, arms slightly spastic, spastic legs in extension, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses. Died 1952. (Probable D.S.)

3.—Serial No. 148, female, born 1924: 1944, aged 20, paræsthesiæ and weak left leg, later paræsthesiæ in fingers, complete remission two months; 1946, loss of vision left eye for one month, paræsthesiæ in fingers and numbness of cheek; 1947, loss of power right leg, which recovered rapidly, but returned later in year; 1948, transient diplopia and incontinence, increasing weakness of legs. Examination 1949, emotional lability, titubation, scanning explosive dysarthria, bilateral pallor both discs. V.A. 6/30, 6/60, nystagmus—fine vertical rotatory in central position and on upward and downward gaze, fast horizontal on lateral gaze, marked tremor on maintaining posture of arms, weak slightly spastic arms, spastic weak legs, generalised hyperreflexia, absent abdominal reflexes and extensor plantar responses Slight wasting of small muscles of both hands. (Probable D.S.)

Brother and sister and her daughter. Brother and sister same environment until 1918.



1.—Serial No. 692, male, born 1896: 1949, aged 53, "cold feeling" from below knees and also in left hand for two months; 1950, increasing ataxia of legs, urgency of micturition and occasional dysarthria; 1951, marked improvement in symptoms. Examination 1951, horizontal nystagmus on left lateral gaze, dysarthria, extensor plantar reflexes. No other abnormal neurological signs. (Probable D.S.)

2.—Serial No. 29, female, born 1914: 1933, aged 19, blind left eye for nine months and paræsthesiæ left leg, with complete recovery. Examination 1935, facile manner, central scotoma left visual field, left abdominal reflexes absent and right sluggish; 1946, slight vertigo. Examination 1946, pallor right disc, nystagmus to right, absent abdominal reflexes, increased knee jerks, doubtful plantar responses; 1948 (no symptoms). Examination, no nystagmus otherwise as in 1946; 1951, no symptoms, examination pallor left disc, nystagmus on right lateral and upward gaze, abdominal reflexes absent, tendon reflexes increased in the right arm and both legs, plantar reflexes extensor, muscle joint sensation defective right great toe. (Probable D.S.)

Brother and sister, same environment.



1.—Serial No. 910, female, born 1918 : 1930, aged 12, diplopia for three weeks; 1948, unsteadiness on walking, diplopia and paræsthesiæ in hands. Examination 1948, euphoria, nystagmus to right and left, slight ataxia of arms, ataxic gait, slightly spastic legs, generalised hyperreflexia, extensor plantar reflexes and bilateral ankle clonus, absent abdominal reflexes. (Probable D.S.)

2.—Serial No. 939, male, born 1920: 1945, aged 25, sudden onset of dysarthria, dyslexia, ataxia and headache (invalided from army with a diagnosis of "acute encephalomyelitis") and when examined in 1947 there were few neurological signs except slight weakness of the handgrips, slightly increased tendon reflexes on the left side, absent abdominal reflexes, extensor plantar reflexes; 1952, pains in the left leg, backache and weakness of left leg for three months. Examination 1952, no abnormal neurological signs except absent abdominal reflexes, increased knee and ankle jerks, plantar reflexes now flexor. (Probable D.S.)

Sister and brother, same address until 1940.



1.—Serial No. 694, male, born 1900: 1938, aged 38, weakness left leg; 1940, both legs weak; 1947, legs weaker, hands weak, precipitancy of micturition, impairment of memory, transient diplopia. Examination 1951, euphoria, horizontal nystagmus to right and left, weak arms, intention tremor right and left, legs weak with slight hypotonia, chairfast, extensor plantar reflexes, absent abdominal reflexes, arm jerks increased, leg jerks not increased, muscle joint and vibration sensation defective in legs. (Probable D.S.)

2.—Serial No. 276, female, born 1907: 1941, aged 34, paræsthesiæ in legs, urgency and incontinence of micturition for three months; 1944, loss of sensation in right arm for two months, followed by loss of sensation left arm for two months; 1945, sudden blindness of left eye, followed by squint and later diplopia for two months, complete recovery except for slight impairment in visual acuity; 1947, gradual onset of numbness of legs and cramps; 1949, sudden blindness left eye for one week. Examination 1950, emotional lability, left-sided temporal pallor. V.A.R. J1, V.A.L. J8, intention tremor right arm, slight weakness right hand grip, weak legs, with spasticity, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses, impaired vibration and muscle joint sensation in legs. (Probable D.S.)

Brother and sister, same environment until 1916.



1.—Serial No. 200, female, born 1897: 1923, aged 26, sudden onset blindness both eyes, paræsthesiæ all limbs and weakness of the legs; in six months made an almost complete recovery except for a little residual weakness of the left leg; 1932, vision again impaired, weakness of legs, especially left, and precipitancy of micturition, and again made nearly complete recovery except for some weakness of the legs; 1939, increasing weakness of legs and impairment of memory, marked precipitancy of micturition; 1942, bedridden. Examination 1950, marked euphoria, dementia, bilateral optic atrophy, with visual impairment, pupils react sluggishly to light, marked ataxia in upper limbs, paraplegia in flexion, generalised hyperreflexia, extensor plantar reflexes, gross impairment of sensation for light touch and vibration in lower limbs. (Probable D.S.)

2.—Serial No. 199, female, born 1900: 1918, aged 18, diplopia for one year; 1930, gradual onset of weakness of legs, with partial recovery over five years. Pregnancies 1936 and 1937, without ill effect. However, following third pregnancy 1939, legs became gradually weaker. Examination 1950, bilateral optic atrophy, visual acuity 6/6 R. 6/6 L., slight intention tremor left arm, spastic ataxia paraplegia, with increased knee and ankle jerks, vibration sense absent in legs, Romberg positive, plantar reflexes extensor. (Probable D.S.)

Both sisters lived at home together until 1935.

(17)

1.—Case H. M., female, born 1890: 1927, aged 37, weakness of legs; 1930, temporary numbness left arm and dysphagia; 1945, weakness left arm. Examination 1947, bedridden, dysarthric, left arm and legs spastic, with limited voluntary movements, ataxic right arm, astereognosis right hand, impossible to test reflexes as legs were markedly contracted. Died 1948. (Possible D.S.)

2.—Serial No. 61, male, born 1908 : 1933, aged 25, precipitancy and frequency micturition; 1935, retention of urine for few days, which recovered, but returned to precipitancy and frequency; 1936, progressive weakness right leg, accompanied by paræsthesiæ; 1947, transient blurred vision. Examination 1947, slight impairment of memory, slight nystagmus on upward gaze, slight dysarthria, spastic in extension, generalised hyperreflexia, abdominal reflexes absent, extensor plantar reflexes. Vibration sense impaired in legs. (Probable D.S.)

Brother and sister, same environment.



1.—Serial No. 487, male, born 1911: 1932, aged 21, gradual increasing weakness of legs. Examination 1950, nil abnormal found in cranial nerves and arms, spastic weakness of legs, increased reflexes, ankle clonus, absent abdominal reflexes, extensor plantar reflexes. (Possible D.S.)

2.—Serial No. 486, female, born 1914: 1939, aged 25, following childbirth, marked impairment of vision for four months; 1949, paræsthesiæ right leg and right foot drop for one year. Examination 1949, temporal pallor right disc, visual acuity normal, weakness right leg especially, dorsiflexors of right foot, right abdominal reflexes absent, impairment of all modalities of sensation up to level of T8. Examination 1950, normal findings except for pallor right disc. (Probable D.S.)

Brother and sister, common environment until sister's marriage in 1937.





1.—Serial No. 436, female, born 1890: 1939, aged 49, weakness and unsteadiness of legs. Examination 1949, euphoria, mild dementia, pallor right disc, with good visual acuity, coarse monocular nystagmus to right and left, weak and ataxic left hand, legs slightly weak, especially left leg, spastic ataxic gait, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses and diminished vibration sensation in legs. (Probable D.S.)

2.—Serial No. 149, male, born 1908: (History not very reliable.) Paræsthesiæ left hand, bilious attacks, recurring diplopia and unsteadiness on feet. Examination 1949, dementia, facile euphoria, coarse monocular nystagmus to right and left, spastic ataxic legs, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, difficulty starting micturition. (Probable D.S.)

Mother and son, same environment.



1.—Serial No. 614, male, born 1884: 1942, aged 58, gradual onset of progressive weakness of legs; 1945, slowness in micturition. Examination 1952, horizontal nystagmus to right and left, spastic legs, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, vibration sensation absent in legs. muscle joint sensation defective in legs. (Investigated fully in hospital in 1945.) (Possible D.S.)

2.—Serial No. 571, male, born 1910: 1945, aged 35, weakness right leg, gradually progressive. Examination 1951, euphoric, weak spastic left arm and leg, left arm jerks increased, leg jerks increased, ankle clonus, left abdominal reflexes absent, extensor plantar reflexes. (Possible D.S.)

Father and son, similar environment.



1.—Serial No. 194, male, born 1916 : 1940, aged 24, numbness right cheek for several days, followed by impairment of vision in right eye and to lesser extent left eye; total recovery in two months. Six months later left leg weak, progressing in a few days to total uselessness lasting for one month; gradual recovery, with slight residual disability; 1942, vision again impaired in both eyes; recovery followed by diplopia lasting two weeks; 1947, difficulty in micturition and defæcation, impotence and increasing weakness of legs. Examination 1949, bilateral optic atrophy, visual acuity R6/18, L6/20, well-marked monocular nystagmus to left and conjugate nystagmus to right. Slight weakness of arms, legs spastic ataxic, with slight weakness, knee and ankle jerks increased, plantar responses extensor, impaired vibration and postural sensation in legs, Romberg's sign positive; 1952, just able to walk. (Probable D.S.)

2.—M. B., female, born 1913: Examination 1946, bedridden, fatuous euphoria, marked nystagmus and optic atrophy. Duration 9-12 years, diagnosed as disseminated sclerosis.

Brother and sister, same environment.

1.—Case T. D., male, born 1896 : 1925, aged 29, sudden paralysis of left arm and leg, left side numb, and diplopia without loss of consciousness. He made a partial recovery, but still has occasional diplopia and stiff legs. Examination 1950, euphoria, fine monocular nystagmus on left lateral gaze, paraplegia in extension, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes. (Possible D.S.)

2.—Serial No. 278, male, born 1925: 1942, aged 17, loss of feeling and control of the right foot and unsteady gait, lasting two months. Six months later clumsiness and loss of feeling in the right hand; 1944, dimness of vision in the right eye for two months; 1947, numbness of the right leg, followed in a few days by "influenza"; when he got out of bed on the seventh day he found he was very unsteady, right leg weak, and had difficulty with micturition. Examination 1947, pallor right disc, fine lateral nystagmus, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses and vibration sensation absent in legs. (Probable D.S.)

Uncle (M) and nephew, no common environment.



(23)

1.—Serial No. 508, male, born 1895: 1935, aged 40, weakness and numbness right arm for one week; 1940, gradual onset weakness left leg and arm especially after exercise, also hesitancy of micturition with "stuttering bladder," both symptoms remain unchanged; 1945, right leg "gave way" and numbness right leg for one week. Examination 1950, euphoria, weakness left arm and leg, hyper-reflexia, absent abdominal reflexes, extensor plantar responses, vibration and muscle joint sensation impaired in legs. (Probable D.S.)

2.—Serial No. 256, male, born 1925: 1948, aged 23, paræsthesiæ and weakness right arm and leg for four weeks; 1949, sudden difficulty speaking and weakness of right arm and leg for a few weeks; 1950, sudden weakness of both legs and dysuria for two months. Examination 1950, slight nystagmus on left lateral gaze, marked impairment of all forms of sensation below level of fourth thoracic segment, with weakness of legs, especially left, absent abdominal reflexes and extensor plantar responses (C.S.F. normal pressure, 4 cells, trace globulin, 45 mgms. per cent. protein, Lange 112200000); 1951, sudden weakness and numbness left arm and leg, precipitancy of micturition and occasional incontinence. Partial recovery; 1952, sudden loss of sight right eye, with recovery in three months. (Probable D.S.)

Father and son, same environment.



1.—Case J. P., female, born 1903: 1950, aged 47, Dr. Stulik, Larchmont, New York, reports that this patient has been ill for twelve years; at present she has dysarthria, ataxia, and weakness of all limbs. W.R. negative. "The diagnosis might be multiple sclerosis."

2.—Serial No. 417, female, born 1911: 1927, aged 16, weakness and ataxia of legs, weakness right hand, diplopia and precipitancy of micturition; after four years nearly complete recovery except for slight weakness right leg; 1946, sudden weakness of legs, with remission for three months, but recurrence and variable but progressive downward course; 1949, blurred vision for two weeks. Examination 1950, temporal pallor left disc, nystagmus on left lateral gaze, intention tremor right arm, legs weak, generalised hyperreflexia, abdominal reflexes absent, plantar responses extensor, vibration sensation diminished in legs; 1951, much improved except for occasional weakness of legs. Examination, mild spastic ataxic gait. (Probable D.S.)

Two sisters lived at home in Ireland together until 1928.



1.—Case G. F., male, died age of 57. Following a fall, he was paralysed for many years before death.

2.—Serial No. 243, male, born 1889: 1935, aged 46, increasing weakness left leg, which spread to the right leg. At times dysarthria, frequency and urgency of micturition. Examination 1950, temporal pallor right disc, slight irregularity of the pupils, slight intention tremor right hand, paraplegia in flexion, impairment of all modalities of sensation in legs, generalised hyperreflexia, absent abdominal reflexes and extensor plantar responses. (Probable D.S.)

3.—D. M. G., male, attended out-patients London Hospital, 1948. Dr. Russell Brain diagnosed disseminated sclerosis. Patient complained that for two years his legs had been weak and his walking unsteady. He also complained of blurred vision and diplopia. There was marked nystagmus to right and left and upwards, slight intention tremor in left arm, the abdominal reflexes diminished, both plantar reflexes were probably extensor, left knee jerk increased, gait spastic ataxic. (Probable D.S.) (26)

1.—Case E. S., female, born 1903, died 1938: Superintendent of mental hospital stated that this was a "typical case of disseminated sclerosis, with emotional outbursts, dysarthria, spastic paraplegia, and absent abdominal reflexes. The C.S.F. showed a marked lange paretic curve with a negative W.R."

2.—Serial No. 73, male, born 1905: 1933, aged 28, sudden ataxia of legs, weakness right leg and "dimness of vision." Six months later marked precipitancy and urgency of micturition, slight deterioration since. Examination 1947, euphoria, slight deterioration in recent memory, temporal pallor right disc, with slight impairment of vision, nystagmus on lateral gaze, dysarthria, tone increased in arms, intention tremor in all limbs, spastic ataxic legs, with slight weakness, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses, Romberg positive. (Possible D.S.)

Brother and sister, similar environment.



1.—Case W. B., male, born 1896, seen by Sir Henry Cohen, 1934, in Liverpool, who commented as follows: "History of paræsthesiæ fourteen years ago, following a fall whilst skating, accompanied by a typical 'barber's chair' phenomenon, affecting the lower limbs and trunk to the waist and also the arms; 1932, cramps in legs and right arm associated with increasing weakness and unsteadiness of gait. Examination showed pale discs, slight nystagmus on extreme lateral gaze, slight weakness and spasticity right arm, spastic legs, generalised hyperreflexia, bilateral extensor toe responses, diminished abdominal reflexes, marked diminution of muscle joint and vibration sensation in legs. I fear there can be no doubt that he is suffering from disseminated sclerosis."

2.—Serial No. 304, female, born 1905: 1928, aged 23, noticed that the thigh muscles were weak after tennis; 1929, "nervous breakdown," couldn't eat, shakiness of legs, paræsthesiæ of head and back for three months; 1931, sudden severe pains in legs, weakness of legs, following influenza, with partial recovery, but had to retire from teaching; 1939, gradually increasing weakness right leg. Examination 1951, right arm paralysed, spastic at shoulder and elbow joints, flaccid at wrist, left arm slight weakness, spastic paraplegia, very limited voluntary movements of legs, chairfast, extensor plantar reflexes, absent abdominal reflexes, Hoffmann reflexes present. All modalities of sensation impaired in legs, postural and vibration sensation impaired in hands. (Probable D.S.)

(28)

1.—Case M. B., female, born 1874, died 1928: Ataxic paraplegia for four years prior to death; had been seen by Professor Sir William Thomson on several occasions.

2.—Serial No. 328, male, born 1902 (the following is a brief résumé of a letter from Dr. Fergus Ferguson, Manchester): 1948, aged 46, paræsthesiæ of legs, increasing weakness of legs; 1950, difficulty in focussing and urgency of micturition. Examination 1950, bilateral pallor of discs, defective movements of eyes to the left, with nystagmoid movements, left plantar reflex indefinitely extensor, right definitely extensor, abdominal reflexes absent, left arm jerks brisker than right, Hoffmann reflexes positive.

3.—Serial No. 115, female, born 1911: 1929, aged 18, dimness of vision (? both eyes) and severe headaches which lasted three months; 1940, severe pains around lower thorax, lasting two months; 1945, paræsthesiæ of legs and progressive ataxia of legs; 1948, frequency and occasional incontinence of micturition and weakness of hands. Examination 1949, euphoria, pallor left disc, V.A.L. 6/6, V.A.R. 6/6, coarse horizontal nystagmus to left, slight weakness of hands, ataxia of arms and legs, increased knee and ankle jerks, absent abdominal reflexes, extensor plantar reflexes, vibration sensation diminished in legs. (Probable D.S.)



1.—Case E. O'R., female, born 1906: 1941, aged 35, numbness and stiffness of hands for several months; 1949, sudden onset of weakness, numbness and stiffness of left leg, followed by similar symptoms in right leg, difficulty in micturition and increasing constipation. Examination 1951 (the Presbyterian Hospital, New York), slight generalised spastic weakness of all limbs, with generalised overactive tendon reflexes, Babinski reflex positive, absent abdominal reflexes, slight ataxia of arms, impaired muscle joint and vibration sensation in legs. All investigations, including myelogram negative. (Diagnosed multiple sclerosis.)

2.—Serial No. 825, female, born 1910: 1950, aged 40, impairment of vision both eyes for six weeks; February, 1951, impairment of vision left eye, followed in two weeks by impairment of vision right eye which lasted two weeks, right leg weak, right hand clumsy; in May, diplopia developed. Examination 1951, euphoria, V.A.L. J14, V.A.R. J8, fundi normal. Complete palsy left VI nerve, horizontal nystagmus on right lateral gaze, slight inco-ordination of arms, spastic ataxic gait, but no weakness, generalised hyperreflexia, absent right abdominal reflexes, extensor plantar reflexes; 1952, much improved, less ataxic and vision normal. (Probable D.S.)

Two sisters, same environment until 1930.

APPENDIX Y.

Case histories of families 30-44 in which we examined one member and the evidence concerning the other was less convincing, such as a letter from the family doctor or a death certificate.



1.—Serial No. 339, male, born 1913: 1937, aged 24, diplopia for three weeks, unsteadiness of gait for one year; 1940, paræsthesiæ of arms and legs and weakness of legs for one month, with complete recovery; 1949, following influenza, weakness, and ataxia of legs for three weeks, with nearly complete recovery; 1950, following sore throat, weak ataxic legs, with increasing disability. Examination 1950, euphoria, bitemporal pallor, V.A.R. and L. 6/6, coarse horizontal nystagmus of monocular type, vertical nystagmus on upward gaze, intention tremor in both arms, slight spasticity right arm, slight spastic ataxia of legs, generalised hyperreflexia, absent abdominal reflexes, doubtful plantar reflexes, stereognosis slightly impaired in hands, vibration sensation absent in legs (Probable D.S.)

2.—Case Wm. C., male, died 1944, aged 30. Death certificate—disseminated sclerosis.

First cousins, different environment.



1.—Female, died aged 38; suffered from disseminated sclerosis for four years.

2.—Male, born 1893, died 1939; paralysed for fourteen years; supposed to have suffered from disseminated sclerosis.

3.—Serial No. 219, female, born 1895: 1924, aged 29, dizziness, blurred vision, weakness of back, partial remission (history not satisfactory). Examination 1951, deteriorated, euphoric, V.A.R. J10, lens opacity left eye, pallor right disc, horizontal monocular nystagmus to right and left, rotatory nystagmus on upward gaze, arms slightly weak and spastic, arms ataxic, especially left, legs paraplegic in flexion, no voluntary movements, generalised hyperreflexia, vibration sensation absent, pain, light touch and muscle joint sensation defective in legs. (Probable D.S.)

Brother, sister, and male cousin. All three cases lived in close proximity.



1.—Serial No. 723, male, born 1887: 1920, aged 33, progressive weakness and unsteadiness of legs; 1946, attack of tic douloureux, with two relapses in period 1946-1951; 1948, sudden weakness left arm. Examination 1951, pallor left disc, with impairment of vision, coarse monocular nystagmus on right lateral gaze, finer monocular nystagmus on left lateral gaze, left facial weakness, slight spastic weakness left arm, spastic paraplegia of legs, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses, vibration and muscle joint sensation absent in legs, light touch and pain sensation impaired below knees. (Probable D.S.)

2.—Case M. H., female, born 1899, and died 1946: Illness started 1941 with "eye trouble and paralysis from the waist down. At the end her speech was affected."



1.—Case M. M., female, died aged 80: Paraplegic for many years.

2.—Case M. E., female, born 1891, died 1951: Twenty years' history of weakness of legs.

3.—Serial No. 207, female, born 1897: 1935, aged 38, retention of urine for four days, followed by difficulty in micturition for three weeks, with full recovery; 1940, gradual onset weakness and stiffness right leg, which progressed. Examination 1951, euphoria, fine nystagmus to right and left, weakness right handgrip, with slight ataxia, spastic paraplegia, generalised hyperreflexia, extensor plantar responses, absent abdominal reflexes. (Probable D.S.)



1.—Case M. W., female, died 1950: Considered by her own doctor to have been a case of disseminated sclerosis of long standing.

2.—Serial No. 833, female, born 1903: 1926, aged 23, vomiting and vertigo for three days, followed by ataxia for ten days; 1932, right leg weak, especially after a long walk, with partial recovery in two months; 1939, gradual increasing weakness of both legs; 1951, left arm weak and unsteady, frequency and precipitancy of micturition. Examination 1951, bedridden, euphoric, coarse horizontal nystagmus, left-sided upper motor neurone facial weakness, ataxic arms, spastic paraplegia in flexion, generalised hyperreflexia, absent abdominal reflexes, extensor plantar responses. (Probable D.S.)



1.—Case A. A. W., female, died 1942, aged 44: Death certificate—disseminated sclerosis.

2.—Serial No. 251, female, born 1903: 1939, aged 36, dragging and weakness right leg, paræsthesiæ right foot, urgency of micturition and increasing ataxia; 1948, difficulty focussing the eyes at intervals. Examination 1950, bilateral optic atrophy, with normal visual acuity, generalised hyperreflexia, absent abdominal reflexes, spastic ataxic weakness of legs, left arm falls away, with pseudo-athetotic movements of fingers, extensor plantar reflexes, impaired muscle joint and vibration sensation in legs and left arm. (Probable D.S.)



1.—Serial No. 899, male, born 1900: 1926, aged 26, diplopia for some months; 1928, sudden loss of sight right eye for four months, with complete recovery; 1931, gradual progressive weakness of legs; 1935, bedridden. Examination 1952, obese, euphoric, pallor left disc, V.A.R. J8, V.A.L. J8, marked monocular nystagmus to right and left and slight vertical nystagmus on upward gaze, no voluntary movements right arm, which is spastic at elbow and flail at wrist, slight weakness left arm, paraplegia in flexion, generalised hyperreflexia, vibration sensation impaired in arms, absent in legs, other forms of sensation slightly impaired. (Probable D.S.)

Case 2.—Case E. S., female: Died, aged 35, from disseminated sclerosis; diagnosis confirmed by Sir Thomas Huston. No notes available.

Brother and sister, lived together until brother went to Canada 1927-28 for a number of years.



1.—Case A. H., female: Died in New York, approximately 1931, from "spinal paralysis."

2.—Case M. H., female, born 1897: Died 1948; had been diagnosed as disseminated sclerosis. Paralysed for seven years prior to death.

3.—Serial No. 395, male, born 1904 (not a reliable witness, on account of mental deterioration): 1946, aged 42, increasing weakness and stiffness left leg. Examination 1949: euphoric, bitemporal pallor, with slight impairment of vision, horizontal monocular nystagmus on lateral gaze, intention tremor in arms, slight weakness and marked spasticity of legs, generalised hyperreflexia, with patellar and ankle clonus on left side, absent abdominal reflexes, extensor plantar responses, diminished vibration sensation in legs. (Probable D.S.)



1.—Serial No. 2, male, born 1900 (history unreliable owing to faulty memory and low intelligence); 1946, aged 46, increasing weakness of legs. Examination 1949, dementia, euphoria, pallor of both discs, V.A.R. 6/9, V.A.L. 6/60, concomitant internal strabismus, horizontal nystagmus to right and left, spastic paraparesis, generalised hyperreflexia, extensor plantar reflexes, absent abdominal reflexes. (Probable D.S.)

2.—Case G. L., female, born 1910: 1943, aged 33, paræsthesiæ in hands and feet and "dizziness"; 1950, attended hospital for "rheumatism"; 1951, admitted to hospital and diagnosed as disseminated sclerosis.



1.—Case E. G., female, born 1879: 1920, aged 41, numbness of left leg, which trailed when she walked; 1922, right leg similarly affected; 1925, difficulty of micturition. Examination 1926 (Royal Victoria Hospital, Belfast; Professor Sir W. W. D. Thomson), nystagmus, spastic weakness of legs, absent abdominal reflexes, increased knee and ankle jerks. Died 1931, diagnosed as disseminated sclerosis.

2.—Serial No. 584, female, born 1918: 1944, aged 26, diplopia and ataxia for three days; 1949, weakness of legs, with partial recovery. Examination 1951, slight spastic weakness of legs, generalised hyperreflexia, absent abdominal reflexes; 1952, sudden increased weakness of legs, with rapid recovery in a few weeks. (Probable D.S.)

Mother and daughter, same environment until 1931.



1.—Case W. S., male, born 1874: Died 1936, paralysed for many years before his death.

2.—Case M. S., female: Died 1943, "paralysed for many years."

3.—Serial No. 586, male, born 1888 : 1935, aged 47, sudden weakness of legs, with pain in back, unable to walk, complete recovery in one month; 1939, blurred vision, especially left eye, for four weeks; 1941, gradual onset of increasing weakness left leg, spreading to right leg. Examination 1951, euphoria, pallor both discs, especially the right, intention tremor right arm, spastic paraplegia, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, slight impairment of muscle joint and vibration sensation in legs. (Probable D.S.)



1.—Case C. B., female, born 1894: Died 1929; "legs became weaker and weaker." Diagnosed as disseminated sclerosis.

2.—Serial No. 192, female, born 1907: 1942, aged 35, numbness and weakness right leg for three months; 1946, gradual progressive weakness right leg, spreading to left leg, diplopia and precipitancy of micturition. Examination 1950, monocular nystagmus on right and left lateral gaze, spastic ataxic legs, generalised hyper-reflexia, absent abdominal reflexes, extensor plantar reflexes, diminished muscle joint and vibration sensation in legs. (Probable D.S.)



1.—Case D. H., male: Died, aged 76; "partially paralysed since age of 30, bedridden for the last four years."

2.—Serial No. 297, male, born 1885: 1920, aged 35, recurring paræsthesiæ left leg and also myoclonic jerks left leg; 1941, gradually increasing weakness of legs. Examination 1949, bilateral optic atrophy. V.A.R. and L. J6, slight weakness left arm, with increased reflexes, weakness and spasticity of legs, with increased tendon reflexes, absent abdominal reflexes, extensor plantar reflexes, absent vibration sensation in legs. (Probable D.S.)



1.—Serial No. 564, female, born 1902: 1939, aged 37, progressive weakness of legs; 1938, gradual onset of impairment of vision of right eye; 1946, "stiffness" of right hand; 1950, retention of urine, following gynæcological operation. Examination 1951, bilateral optic atrophy, V.A.L. and R. J11, horizontal nystagmus on lateral gaze, monocular to right, intention tremor in arms, slight weakness in arms, spastic paraparesis, generalised hyperreflexia, absent abdominal reflexes, extensor plantar reflexes, vibration sensation absent in legs, muscle joint sensation impaired in toes, slight hypalgesia and hypæsthesia right arm and leg, two-point discrimination impaired right hand. (Probable D.S.)

2.—Case M. C., female, born 1903: Died 1945, diagnosed disseminated sclerosis at Mater Hospital, Belfast (no notes available).



1.—Case B. McR., female : Dead, diagnosed as disseminated sclerosis; paralysed for fifteen years. Doctor writes—"Bedridden, euphoric, dysarthric, intention tremor of arms. I have no doubt that she suffered from disseminated sclerosis."

2.—Serial No. 388, female, born 1912: 1949, aged 37, numbness of scalp which lasted three weeks, followed by sudden blindness of right eye, which made a partial recovery; 1950, "dizzy attacks," paræsthesiæ of right leg, followed by paræsthesiæ of left arm, pains in legs, and weakness of ankles, with partial recovery; 1952, increased weakness of legs, with increasing pains in back for three months, with partial recovery. Examination 1952, pallor right disc, with V.A.R. J12, slight weakness of legs, increased knee and ankle jerks, extensor plantar reflexes. (Probable D.S.)

Niece and maternal aunt, same town until 1946.

APPENDIX Z.

THER	NEUROLOGICAL	DISORDERS	AMONG	тне]	Near	RELATIVES.
	THER	THER NEUROLOGICAL	THER NEUROLOGICAL DISORDERS	THER NEUROLOGICAL DISORDERS AMONG	ther Neurological Disorders among the 1	THER NEUROLOGICAL DISORDERS AMONG THE NEAR

Chronic neurological	disorders	s wit	hout a di	agnosis	-	-	18
Psychosis -	-	-	-	-	-	-	16
Mentally defective	-	-	-	-	-	-	7
Psychoneurosis	-	-	-	-	-	-	5
Parkinson's disease	-	-	-	÷	-	-	5
Muscular dystrophy	-	-	-	-	-	-	2
Epilepsy -	-	-	-	-	-	-	1
Vascular accidents	-	-	-	-	-	-	1
Subacute combined d	egenerati	on of	the cord	-	-	-	1
Myasthenia gravis	-	-	-	-	-	-	1

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