

The Condition of the Peripheral Nerve in Leprosy Under Various Forms of Treatment

Conduction Velocity Studies in Long-Term Follow-Up^{1, 2}

Alexander Magora, Jacob Sheskin, Felix Sagher and Benjamin Conen³

Peripheral nerve lesions are common in leprosy. They may be of two basic types: infiltrative, with inflammation and fibrosis along the nerve, or constrictive, in a patchy form, with signs of compression or strangulation. The nerve lesions may have an insidious, gradual onset with mild clinical signs or they may occur suddenly, especially during leprosy reactions. In many instances therapy may arrest the progress of the nerve damage.

The question arises, however, as to whether, in the absence of leprosy reaction (LR), the fibrosis of the nerve does not sometimes cause an insidiously progressive lesion, without immediately apparent clinical symptoms or signs. An additional problem to be considered is the type of therapy which most effectively influences the nerve lesion.

The sulfones (derivates of 4,4'-diaminodiphenyl sulfone) were first introduced in 1937 by Buttle *et al.* (⁷) and Fourneau *et al.* (¹⁵) against streptococcal infections, and later by Rist *et al.* (³⁷) and Feldman *et al.* (¹³) against animal tuberculosis. In 1941, after further assessment (¹⁰) they were tried in human leprosy (¹²). The good results obtained warranted their wide acceptance as the most commonly used

therapeutic agent in leprosy. It should, however, be stressed that among other toxic effects, the sulfones may cause mild neuritis, manifested by paresthesias and dysesthesias (⁴³).

A second group of drugs, mostly used in recalcitrant cases of LR, are the steroids, mainly prednisone and triamcinolone. These drugs may cause, in addition to their metabolic effects, obvious histologic, electromyographic and clinical signs of myopathy or polymyositis. This has been experimentally demonstrated in animals (^{3, 11, 25}) and has been observed in human subjects suffering from various clinical conditions (^{5, 8, 20, 35}). It would also seem that the addition of fluorine in position 9 (as in triamcinolone) further enhances the appearance of steroid myopathy (^{2, 26, 31, 44}).

In 1965, thalidomide [alpha- (N-phthalimido)-glutarimide or N-phthaloyl-glutarimide], first investigated by Kunz *et al.* (²⁹) and previously used as a powerful, nondepressant hypnotic, was introduced as a therapeutic agent with dramatic effects on the neuritis encountered in the leprosy reaction of the lepromatous type (^{40, 41}). However, notwithstanding its influence on the polyneuritis of leprosy and the lack of neurotoxic effects in animals (²⁸), thalidomide has been incriminated as the cause of polyneuropathy in human beings, especially if administered for prolonged periods of time (^{9, 14, 16, 17, 21, 38}).

The question therefore arises in any long-term follow-up of leprosy, as to whether the condition of the peripheral nerve deteriorates, with or without LR and, if it does, whether the additional pathologic changes are primarily caused by the basic

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³ A. Magora, M.D., Department of Physical Medicine and Rehabilitation; J. Sheskin M.D., Department of Dermatology and Venereology; F. Sagher, M.D., Department of Dermatology and Venereology; and B. Conen, M. Sc., Electronics Engineer, Electrodiagnostic Research Unit, Hadassah University Hospital & Government Hospital for Hansen's Disease, Jerusalem, Israel.

disease, by the various drugs employed or by a combined effect of both factors.

The parameter selected for the investigation of the nerves in leprosy is the conduction velocity of the motor fibers (MCV), a test whose value and reliability in a variety of clinical conditions involving the peripheral nerve has been proven (18, 19, 33, 42).

In the following report we present the results of a six-year follow-up study of leprosy patients who received various therapeutic regimes.

MATERIALS AND METHODS

The investigation was carried out on 103 leprosy patients, of whom 61 were men and 42 women. Sixty-seven were of the lepromatous, eight of the tuberculoid and 20 of the indeterminate type; eight had the dimorphous (intermediate) form.

The classification was based on the clinical symptomatology, biopsy findings, bacteriologic examination and lepromin test. No clear cut changes from one type to another have been observed during this period.

With regard to the cases of "indeterminate type" they could certainly have changed into lepromatous or tuberculoid types provided they remained without therapy. All these patients were, however, treated with sulfones, as a result of which their symptoms have slowly disappeared.

These 103 leprosy patients were followed for a period of six years. After the initial evaluation they underwent repeated medi-

cal and bacteriologic examinations at 6 - 12 month intervals. The medical investigation consisted of a general, dermatologic and neurologic examination; the latter included examination of the muscle power, sensory status, tendon and periosteal reflexes and the presence of Tinel's sign, atrophy or nerve thickening.

The relationship between the type of disease and age appears in Table 1 and between type of disease and duration since onset in Table 2. Table 3 shows the relationship of the disease to the clinical and bacteriologic status. Fifty-two cases were in an arrested state, and all were negative bacteriologically. Fifty-one were clinically active and of these, 26 were bacteriologically negative and 25 positive.

The patients received three types of treatment: sulfones alone or sulfones in combination with either prednisone or thalidomide. Table 4 presents the various therapeutic combinations according to the type of disease. All patients received sulfones; 15 received steroids (prednisone) and 27 thalidomide in addition to the sulfones. Prednisone and thalidomide were never administered together, but some patients who had repeated LR, received either prednisone or thalidomide in their recurrent attacks. The basic indication for the use of either prednisone or thalidomide was the presence of a LR; in some instances, as will be seen, the patient continued with the same combination of sulfone and either prednisone or thalidomide for longer periods. The usual dose of sulfone

TABLE 1. Relationship between type and form of leprosy and age of patient.

Age (yrs.)	Type				Total
	Lepromatous	Tuberculoid	Indeterminate	Dimorphous	
Up to 20	1	—	—	—	1
21-30	13	1	6	0	20
31-40	15	2	5	1	23
41-50	10	1	2	2	15
51-60	18	2	2	3	25
61 & over	10	2	5	2	19
Total	67	8	20	8	103

TABLE 2. Relationship between type and form of leprosy and time interval after onset of disease.

Onset of disease (yrs.)	Type				Total
	Lepromatous	Tuberculoid	Indeterminate	Dimorphous	
1-5	1	—	—	1	2
6-10	13	—	3	1	17
11-15	13	—	2	2	17
16-20	12	5	8	—	25
21-25	8	3	4	4	19
26-30	5	—	—	—	5
31 & over	15	—	3	—	18
Total	67	8	20	8	103

was 100 mgm. which was gradually decreased to an average of 25 mgm. daily. The daily dose of prednisone was 5-30 mgm. and of thalidomide 300-500 mgm. initially, with a gradual decrease to a maintenance dose of 50-100 mgm./day.

Of the 103 patients, 38 had a total of 127 LR during the six-year follow-up period. Most of the LR were spontaneous but some of them were provoked by discontinuation of the sulfone therapy. The type and duration of the disease, and the age and sex of the patients who had LR are presented in

Table 5. The clinical manifestations and the clinical and bacteriologic status appear in Table 6. From these two tables it can be seen that the most common manifestations were erythema nodosum and neuritis, either separately or together. The clinical manifestations of recurrent LR in the same subject were similar. Of the 38 patients, 12 were clinically in an arrested state and 11 of them were negative bacteriologically. Twenty-six were clinically active and these were all positive bacteriologically. The general data for the 28 patients in whom

TABLE 3. Relationship between type and form of disease and clinical activity and bacteriologic status.

Clinical status	Bacteriologic status	Type				Total
		Lepromatous	Tuberculoid	Indeterminate	Dimorphous	
Arrested		25	8	18	1	
	Negative	25	8	18	1	52
	Positive	—	—	—	—	—
Active		42	—	2	7	
	Negative	19	—	2	5	26
	Positive	23	—	—	2	25
Total		67	8	20	8	103

neuritis was the prominent manifestation of LR appear in Table 7 and the clinical condition and bacteriologic status in Table 8. From these tables it is clear that none of these factors (type of disease, age, sex, period since onset of disease, clinical condition and bacteriologic status) bear any relation to the appearance or development of neuritis as part of the LR.

As already mentioned, either prednisone or thalidomide were administered, in addition to sulfone, at the time of appearance of the LR, sometimes for prolonged periods. Tables 9 and 10 present the accumulated periods of treatment with prednisone and thalidomide respectively, over the six year period of follow-up. Nine patients received prednisone and 21 thalidomide for more than four months, 18 received thalidomide

for more than one year and 10 for more than three years. From Table 11, in which only the longest, uninterrupted period of therapy is considered, it can be seen that prednisone was most commonly administered for one to six months while thalidomide was used for more than five months in 15 patients and for periods of more than three years in three patients.

Each patient underwent, in addition to medical and bacteriologic examination, an investigation of the motor conduction velocity (MCV) at the time of each visit so that there were at least five follow-up tests per patient. In addition, in the patients with LR, the MCV was examined at least three additional times if the manifestations were dermatologic only, and daily or at two-day intervals if neuritis was the promi-

TABLE 4. Relationship between type of disease and therapy.

Therapy	Type				Total
	Lepromatous	Tuberculoid	Indeterminate	Dimorphous	
S ^a	42	8	17	4	71
S + P ^b	4	—	1	—	5
S + T ^c	13	—	1	3	17
S + P + T	8	—	1	1	10
Total	67	8	20	8	103

^a S = Sulfone.

^b P = Prednisone.

^c T = Thalidomide.

TABLE 5. Type of disease, age, sex and time interval after onset of disease in patients with leprosy reactions.

Type of disease	No. pts.	Age (yrs.)	No. pts.	Sex	No. pts.	Period since onset of disease (yrs.)	No. pts.		
LL	31	21-30	9	Male	21	1-5	1		
LT	1	31-40	8	Female	17	6-10	7		
LI	2	41-50	4			11-15	7		
LD	4	51-60	11			16-20	6		
		61 & over	6			21-25	8		
						26 & over	9		
Total LR	38		38		38		38		

TABLE 6. *Clinical manifestations and condition and bacteriologic status in patients with leprosy reactions.*

Clinical manifestations of LR	No. pts.	Clinical condition	No. pts.	Bacteriologic status	No. pts.
Erythema nodosum	18	Arrested	12	Negative	11
Neuritis	28	Active	26	Positive	27
Other ^a	2				
Erythema nodosum and neuritis	19				
Total LR	38		38		38

^a Iritis, epididymitis.TABLE 7. *Type of disease, age, sex and time interval after onset of disease in patients with leprosy reaction manifested by neuritis.*

Type of disease	No. pts.	Age (yrs.)	No. pts.	Sex	No. pts.	Period since onset of disease (yrs.)	No. pts.
LL	25	21-30	8	Male	17	1-5	0
LT	1	31-40	6	Female	11	6-10	6
LI	1	41-50	2			11-15	7
LD	1	51-60	8			16-20	6
		61 & over	4			21-25	4
						26 & over	5
Total LR	28		28		28		28

nent sign. In most cases the MCV was also examined 3-5 days and 10-15 days after the signs of LR had subsided.

The MCV investigations were carried out with a constant voltage stimulator, up to 200 V., duration of pulse one millisecond (m. sec.) and frequency of one pulse per second. The stimulator had very good ground isolation. The stimulus was given by a two-pronged stimulator and the responses were recorded through a teflon coated concentric needle inserted into the most distal muscle. The response was amplified from 400 to 1,000 times; it was recorded on tape and later accumulated and measured by a computer of average transients (CAT). The CAT was triggered by the stimulus artefact and the responses were synchronously summated in the CAT memory. The results were obtained in analogue form, displayed on an oscilloscope

and were then either photographed or plotted on ultraviolet paper. The advantages of this technic are the accuracy of measurement of the period of latency, the duration of the evoked response, and the control over the consistency of response. An additional advantage is that processing and

TABLE 8. *Clinical condition and bacteriologic status in patients with leprosy reaction manifested by neuritis.*

Clinical condition	No. pts.	Bacteriologic status	No. pts.
Arrested	7	Negative	6
Active	21	Positive	22
Total LR	28		28

TABLE 9. Relationship of prednisone therapy to type of disease and period of administration.

Period ^a	Type				Total
	Lepromatous	Tuberculoid	Indeterminate	Dimorphous	
15-30 days	2	—	—	—	2
1-2 months	1	—	1	—	2
3-4 "	1	—	1	—	2
5-6 "	4	—	—	1	5
7-9 "	2	—	—	—	2
10-12 "	1	—	—	—	1
1-2 years	1	—	—	—	1
Total	12	—	2	1	15

^a Not continuously—in accumulated periods.

measurement can be undertaken after the patient has been discharged and that the procedure is not time consuming. In cases in which the evoked response was of very low amplitude, or where it tended to be obscured by either spontaneous activity or lack of complete relaxation, the CAT was able to extract the real evoked response by exact, synchronous accumulation. The MCV examinations were carried out in a room with constant temperature, always using the same technic. As it was found that the most commonly damaged nerve was the ulnar, all MCV tests were carried out on this nerve, on both sides in each patient. In patients with LR, only the most

severely damaged ulnar nerve was followed up. Before the actual examination, the procedure was explained to the patient and a brief period was provided for relaxation. The ulnar nerve was examined from at least three points: axillary, above the elbow and at the level of the wrist.

Only supramaximal current was used. The distance was measured with an elastic tape, with the elbow in full extension and the wrist in the neutral position, special care being paid not to pull the skin. For MCV of the ulnar nerve the recording electrode was inserted in the abductor digiti minimi muscle. In a number of cases, depending on the clinical condition, the

TABLE 10. Relationship of thalidomide therapy to type of disease and period of administration.

Period ^a	Type				Total
	Lepromatous	Tuberculoid	Indeterminate	Dimorphous	
15-30 days	2	—	—	1	3
1-3 months	2	—	1	—	3
4-6 "	5	—	—	—	5
7-9 "	—	—	—	1	1
10-12 "	—	—	—	—	—
1-2 years	7	—	—	1	8
3 or more years	8	—	1	1	10
Total	24	—	2	4	30

^a Not continuously—in accumulated periods.

TABLE 11. Longest period of continuous administration of prednisone and thalidomide.

Longest period	Prednisone	Thalidomide
Less than one month	2	2
1-2 months	4	6
3-4 "	4	4
5-6 "	3	1
7-9 "	1	1
10-12 "	—	3
1-2 years	1	7
3 or more years	—	3
Total	15	27

median, radial and common peroneal nerves were also examined (Table 12).

The interpretation of the MCV was based on the analysis of three basic criteria: velocity, latency, and duration and form of the evoked response. The velocity was con-

sidered normal if the results obtained were between 50 - 69 m/sec. for the ulnar nerve, 47 - 66 for the median, 50 - 72 for the radial and 43 - 57 for the common peroneal nerve (^{23, 24, 34, 39}). The latency was interpreted according to the segment of nerve examined and the distance to the recording electrode. The age of the patient and variations of temperature (either in the laboratory or of the patient) were also considered. The final interpretation of MCV was also based on comparison of values received on both sides: a difference in velocity of more than 10 m/sec., even if both results were within normal limits, was considered abnormal for the slower conducting nerve. Since a tendency to slower conduction on the nondominant side has been observed (^{4, 30}), the handedness of the patient was also taken into account. In order to avoid any diurnal variations of MCV (²²), most investigations were carried out at approximately the same time of day.

TABLE 12. Results of initial examination of MCV^a in 103 patients.

Nerve	Velocity		Distal latency		Duration of evoked response		Total Examined
	Normal	Abnormal	Normal	Abnormal	Normal	Abnormal	
Ulnar No. %	156 75.7	50 24.3	151 73.3	55 26.7	150 72.8	56 27.2	206 100.0
Median No. %	68 87.0	9 13.0	66 84.4	11 15.6	66 84.4	11 15.6	77 100.0
Radial No. %	43 93.5	3 6.5	43 93.5	3 6.5	43 93.5	3 6.5	46 100.0
Common peroneal No. %	91 84.2	17 15.8	90 83.3	18 16.7	89 82.4	19 17.6	108 100.0
Total No. %	358 81.9	79 18.1	350 80.0	87 20.0	348 79.6	89 20.4	437 100.0

^a The listing is in terms of the number of nerves examined.

RESULTS

The 103 leprosy patients underwent, during the six-year period of follow-up, a total of 1,153 MCV examinations. Of these 437 were carried out at the time of the initial examination, 308 in the follow-up of the 65 patients without signs of LR and 408 in those with various signs of LR (Tables 5-8). The results of MCV were divided into two groups—normal and abnormal.

The results of the initial examination are presented in Table 12 in which they are related to individual nerves. The highest percentage of abnormal MCV was obtained from the ulnar (24.3%) and the lowest from the radial nerve (5.61%). In a few cases, while the velocity was within normal limits, both the latency and the duration of the evoked response were abnormal, indicating a lesion of the distal part of the nerve.

TABLE 13. *Results of MCV after six years follow-up in 65 patients without leprosy reaction.*

Nerve	Normal	Ab-normal ^a	Total
Ulnar No. %	112 ^b 68.9	18 31.1	130 100.0
Median No. %	25 84.2	5 15.8	30 100.0
Radial No. %	14 93.3	1 6.7	15 100.0
Common peroneal No. %	54 ^c 79.4	14 20.6	68 100.0
Total No. %	309 75.4	101 14.6	410 100.0

^a According to latency, velocity and/or duration of evoked response.

^b Of 121 ulnar nerves previously normal.

^c Of 58 common peroneal nerves previously normal.

Table 13 presents the results of MCV in the 65 patients without leprosy reaction, at the end of the six-year follow-up period. A certain deterioration of the MCV, most prominent in the ulnar nerve, may be observed. While 121 ulnar nerves in these patients were conducting normally at the beginning of the follow-up period, only 112 were doing so at the end of this period. The same deterioration, but to a lesser degree, was noted in the common peroneal nerve. This decrease of MCV was never abrupt; repeated examinations showed a gradual slowing down towards abnormal values, accompanied by an insidious clinical loss of motor power. In most instances the duration of the evoked response was the first parameter to change. Throughout the follow-up period these patients received only sulfone and their clinical condition and bacteriologic status did not show any fluctuation.

The situation was completely different in the 38 patients who had had at least one leprosy reaction (LR) and especially so in those with a resultant neuritis. Tables 14-16, in which various relations are presented, refer only to the number of patients or of LR and not to the total number of nerves examined. The reason for this is that in many cases only one nerve, in general the most severely damaged (ulnar) was examined repeatedly, mainly because of pain and the apprehension of the patient. The clinical diagnosis of neuritis as the manifestation of LR was based on the appearance of severe pain in the distribution of at least one peripheral nerve, and possibly thickening of the nerve with or without a positive Tinel's sign, dysesthesia, loss of sensation and motor deficit.

Table 14 presents the results of MCV in relation to the number of days since the appearance of clinical signs of LR treated by sulfones only. The value of "normal" CV was taken to be that found constantly in the previous examinations of the same nerve, regardless of whether this was within the usually accepted normal values or not. Only those values which, on comparison with the previous ones, showed a velocity slower by at least 10 m/sec. and/or a prolonged latency period were regarded as

TABLE 14. Results of MCV in relation to the time interval after onset of leprosy reaction—without antireactional treatment.

MCV	Period in days				
	1-2	3-4	5-7	8-10	11-14
Normal ^a	27	9	8	7	7
Abnormal ^b	11	29	30	31	31
Total	38	38	38	38	38

^a Previous MCV values considered as normal.^b Abnormal values considered only if MCV decreased by at least 10 m/sec.

abnormal. Table 14 shows that of the 38 patients, 11 had an abnormal CV at the end of the second day after the onset of clinical signs, 29 within four days, 30 at the end of one week and 31 after eight days or more. A few remarkable facts emerge from the analysis of this group which, not having received any antireactional therapy, may be considered as a control group. First, in the majority of cases the MCV became abnormal within four days after the onset of the LR. Second, of the 10 patients who had erythema nodosum as their only clinical manifestation of LR, without any clinical evidence of nerve involvement, seven had a constantly normal MCV. On the other hand, in the remaining three patients who started their LR with erythema nodosum only and exhibited an abnormal MCV by the end of one week, clinical evidence of nerve lesions developed in the following days. This finding will be discussed later. A third point of interest, not reflected in this table, is that in the great majority of LR the

ulnar nerve was the one most frequently damaged, in general slightly more so on one side, but without relation to age, handedness or previous degree of damage. The common peroneal nerve was next in frequency, followed by the median. The radial nerve was only very rarely damaged, either clinically or electrically, in the present series of patients.

Of the 89 LR in the 28 patients with neuritis as the major clinical manifestation, 63 were treated with thalidomide and 26 with prednisone. In recurrent LR these therapeutic trials were often carried out consecutively in the same patient. Antireactional therapy was instituted within two days of the onset of LR.

Table 15 shows the results of MCV in the 26 LR treated with sulfone and prednisone. The incidence of nerve lesions, as manifested by abnormal MCV and parallel clinical signs, increased within the first week of the onset of LR. It was only after the first week and especially during the

TABLE 15. Results of MCV in relation to the time interval after onset of lepra reaction treated with prednisone.

MCV	Period in days						
	1-2	3-4	5-7	8-10	11-14	15-21	22-30
Normal	10	5	4	7	10	14	19
Abnormal	16	21	22	19	16	12	7
Total	26	26	26	26	26	26	26

TABLE 16. Results of MCV in relation to the time interval after onset of lepra reaction treated with thalidomide.

MCV	Period in days						
	1-2	3-4	5-7	6-10	11-14	15-21	22-30
Normal	19	18	29	46	59	61	61
Abnormal	44	45	34	17	4	2	2
Total	63	63	63	63	63	63	63

third and fourth weeks that the nerves returned to their previous state. It must, however, be stressed that in seven patients the nerve examined failed to return to its previous state in spite of relatively high dosages (30 mgm./day) of prednisone. The typical neuralgic pain tended to lessen and to disappear gradually after four to seven days of prednisone therapy.

Table 16 shows the MCV in 63 LR treated with sulfone and thalidomide. It is obvious that after only four days the number of nerves affected started to decrease, and within two weeks the great majority of those examined had returned to their previous condition. The neuralgic pain tended to disappear two to five days after the thalidomide therapy was initiated. Only two patients failed to fully respond to this treatment.

Five patients received thalidomide for periods of one month to four years, without suffering from LR. None of these patients developed any signs of LR during this period. Furthermore, 10 patients who had had at least one LR, continued the thalidomide therapy for periods of at least one year. During the thalidomide administration none of these patients suffered from a LR.

Of the 27 patients treated repeatedly with thalidomide, four complained of paresthesias and dysesthesias of the extremities which, after brief periods of time, disappeared spontaneously. It was not certain whether these symptoms had been present before the administration of thalidomide. In any case, it should be stressed that continued thalidomide therapy neither aggravated these complaints nor caused

any other untoward effect. In a fifth patient, after three weeks of thalidomide therapy and while the signs of a LR neuritis were receding, a widespread polyneuritis occurred, with severe neuralgic pain, of the hands and feet, sensory loss and motor deficit of most of the limb muscles, more pronounced distally. A lumbar puncture carried out at this time showed a marked rise of the proteins in the cerebrospinal fluid without increase of cells; a diagnosis of Guillain-Barré syndrome was therefore indicated. The thalidomide was discontinued. Within three months all neurologic signs nearly disappeared. Thalidomide therapy was reinstated and the patient, who at the present writing has received doses of 100 mgm. daily for over one year, does not have any sign of LR or neuritis. It should also be stressed that none of the four previously mentioned patients suffered from any recurrence of dysesthesias or paresthesias, or any other neurologic sign, in spite of administration of thalidomide for prolonged (over one year) periods of time.

None of the 15 patients who received prednisone developed signs of either neuropathy or myopathy. On the other hand, some of those who received the drug for periods of over 6 months developed diabetes, and signs related to sodium retention.

No relationship was found between the effect of thalidomide and prednisone and the type or form of leprosy, although the number of tuberculoid and dimorphous patients is admittedly too small for valid, significant conclusions. No relationship was found with regard to the age of the patient, the time interval since the onset of the disease, clinical activity, bacteriologic

status or the number of nerves damaged. It should, however, be stressed that while both antireactional drugs influenced the course of LR, thalidomide had a dramatic effect, especially when neuritis was the prominent manifestation.

DISCUSSION AND CONCLUSIONS

A number of important conclusions can be drawn from this investigation, but at the same time several questions and problems are raised.

As already demonstrated in a previous study (²²), it is apparent that leprosy patients may, while under sulfone therapy and without LR, show a gradual, insidious deterioration of the condition of the peripheral nerve. This finding probably indicates that the peripheral nerve is subjected to increasing pressure with resultant hypoxia from the presence of even a few adhesions. The MCV tests show that this damage may occur either in the trunk or distal branches, and that it is most common in the ulnar nerve.

With regard to LR, it is obvious that sulfones have no beneficial effect whatsoever upon its course and that without antireactional therapy, the nerve lesion progresses as if no therapy was being provided. On the other hand, both prednisone and thalidomide had a marked effect upon the LR. The effect of prednisone is slower, and that of thalidomide faster and often dramatic when nerve lesions are present. The mechanism of action of these two drugs is still unexplained. While it may be assumed that the antiphlogistic, analgesic, fibrinolytic and anti-immune effects of prednisone play a role, it is so far unknown whether thalidomide possesses such modes of action.

In this connection, it should also be mentioned that while LR occurred under combined therapy with sulfone and prednisone, it did not do so when thalidomide was being used. Although the number of patients is too small for evaluation of statistical significance, it is remarkable that the few patients who had suffered from numerous repeated LR (with lepromatous leprosy) in the past, had no new LR for periods of over three years while under thalidomide treatment.

The problem of thalidomide polyneuropathy also raises some questions. Although reports were found in the literature that thalidomide may cause nerve lesions (^{9, 14, 16, 17, 21, 38}), it is very difficult to understand the mechanism of this process. It has been stressed that the thalidomide polyneuropathy usually occurs after a minimum of three to six months of administration, often after 18 to 24 months (^{6, 14, 17, 36}) and only after the intake of large amounts (¹⁷). If a direct toxic action is considered likely, the neuropathic effect of thalidomide should have occurred more frequently in leprosy patients, in whom the nerves may already be partially damaged and who receive the drug for long periods and in large doses. Of the five patients in our series who developed additional signs of polyneuropathy, one almost certainly had a Guillain-Barre syndrome while in the remaining four the signs were probably caused by the underlying disease. This opinion is strengthened by a number of facts. First, the symptoms of neuropathy were observed in these patients after only three to five weeks of thalidomide therapy and after the intake of relatively small quantities of the drug; furthermore it is not sure whether these complaints were not present before the administration of the drug. Second, the signs and symptoms disappeared completely within a few weeks or months; this is in contrast to the descriptions of thalidomide neuropathy in the literature (^{17, 21}) which stress that in most cases, even partial amelioration may take years after the discontinuation of the drug. Third, the same five patients who developed signs of neuropathy again received thalidomide after the neurologic signs had receded, sometimes for periods of more than one year, and yet none of them developed further signs of neuropathy. The few histologic studies carried out in thalidomide polyneuropathy (^{17, 27}) have reported degeneration of axis cylinders and of the myelin sheath, without segmental demyelination, and a selective loss of nerve fibers with large diameter; these findings indicate only that a neuropathy is present, without, however, being specific for any particular type (¹).

We have tentatively concluded, therefore, that at least in our patients, no neurologic signs related to thalidomide were observed. However, even if mild paresthesias and dysesthesias were to occur because of thalidomide, it is our impression that this should not constitute a contraindication to the use of the drug, because of the excellent effect it has on LR and because all other drugs presently used in leprosy and LR have similar or worse side effects. Our opinion, based on the reported experience with the drug, is that until absolute objective and well controlled proof that thalidomide has toxic effects upon the peripheral nerve is presented, the drug should be further used in leprosy.

The MCV has been proven to be a good, reliable tool for following the condition of the nerve trunk in leprosy, and it is our impression that it is an accurate indicator of the effect of any drug used. While it has on one hand, the disadvantages of being influenced by many intrinsic and extraneous factors and of having wide normal limits, it is, on the other hand, almost the only tool that allows comparative studies of the nerve trunk. This is important in leprosy, and especially in LR, because the main clinical manifestations are often related to the nerve trunk alone, and may sometimes be only subjective.

In the present investigation of LR, the MCV showed decreased velocity and increased latency within 48 hours of the appearance of pain, often before nerve thickening was apparent. As such, it is of value in the objective determination of the presence of LR neuritis. Moreover, while in some cases MCV showed only a mild decrease in velocity, in others it was unrecordable or of extremely low velocity. If many more cases are studied, and the findings correlated with sensory conduction and histologic studies, it should be possible to quantitate, to a certain extent, the degree of nerve damage. MCV studies have been shown to be of value not only in patients in whom neuritis was evident clinically, but also in a few instances in which the dermatologic lesions were predominant. A slowing of velocity, prolonged latency and increased duration of the evoked response,

indicated the possibility of a subclinical neuritis, which was proven by the appearance of clinical signs a few days later.

The fact that the nerve may undergo progressive constriction, can be demonstrated only with the help of electrical tests and, if the changes are in the nerve trunk, by MCV. Clinical examination cannot distinguish mild changes in the sensory or motor status, and the patient may even be unaware of them because of their insidiousness and his own ability to adapt to the changes produced.

The significance of MCV as a prognostic tool was demonstrated by the degree of improvement on treatment which, except for pain, always preceded any clinical improvement by two to five days. This is important in assessing the value of the drug used, in deciding whether the most effective drug has been administered and in determining the degree of residual damage after leprosy reaction.

Finally, it should again be stressed that regardless of its many advantages, the MCV is useful only if the test is carried out under standardized conditions and if provisions are made for the elimination of as many causes of error and artifacts as possible.

SUMMARY

One hundred and three leprosy patients were followed for a period of six years. During this time they underwent regular clinical, bacteriologic and motor conduction examinations.

Of the 103 patients, 38 had 127 leprosy reactions, 89 of them with neuritis. Sixty-three of the latter were treated with thalidomide and 26 with prednisone. These two antireactional drugs were evaluated with regard to their effect upon the neuritis (by clinical and electrical means) and their possible toxic effects.

In some of the patients without leprosy reaction the MCV demonstrated a slow deterioration of the nerve trunk. In a few cases of leprosy reactions manifested by erythema nodosum, the presence of subclinical neuritis was demonstrable.

All the means of evaluation showed thalidomide to have a more rapid effect than prednisone. Thalidomide, in low therapeutic

tic doses for long periods, also apparently succeeded in preventing the recurrence of leprosy reactions. The complications observed in five patients could not be related to thalidomide and were probably caused by the underlying disease in four instances, and by a Guillain-Barre syndrome in the fifth case.

The significance and possible pitfalls of MCV are discussed.

RESUMEN

Se estudiaron 103 pacientes de lepra durante un periodo de seis años. Durante este tiempo se les practicaron regularmente exámenes clínicos, bacteriológicos y de conducción motora.

De los 103 pacientes, 38 tuvieron 127 reacciones leprosas, 89 de ellas con neuritis. Sesenta y tres de estas últimas se trataron con talidomida y 26 con prednisona. Estas dos drogas antireaccionales fueron evaluadas con respecto a su efecto sobre la neuritis (por medios clínicos y eléctricos) y sus posibles efectos tóxicos.

En algunos de los pacientes sin reacción leprosa la VCM demostró un lento deterioro del tronco nervioso. En unos pocos casos de reacciones leprosas manifestadas por eritema nodoso, se demostró la presencia de neuritis subclínica.

Todos los medios de evaluación mostraron que la talidomida tiene un efecto más rápido que la prednisona. La talidomida, en dosis terapéuticas bajas, utilizada durante largos periodos, también logró aparentemente prevenir la recurrencia de reacciones leprosas. Las complicaciones que se observaron en cinco pacientes no se pudieron relacionar con la talidomida y fueron causadas probablemente por la enfermedad subyacente en cuatro de los casos y por un síndrome de Guillain-Barré en el quinto.

Se discute el significado y posibles errores del VCM.

RÉSUMÉ

Cent et trois malades souffrant de lèpre ont été suivis pour une période de 6 ans. Au cours de ce temps, ils ont été soumis à des examens réguliers portant sur la clinique, la bactériologie et la conduction motrice.

De ces 103 malades, 38 ont présenté au total 127 réactions lépreuses, 89 de celles-ci étant accompagnées de névrite. Soixante-trois de ces dernières complications névritiques ont été traitées par la thalidomide, et 26 par la prednisonne. Ces deux médicaments anti-réactionnels ont été évalués en ce qui concerne

leur action sur la névrite par des méthodes cliniques et électriques; leur action toxique possible a également été étudiée.

Chez quelques uns des malades ne souffrant pas de réaction lépreuse, le MCV a révélé une détérioration lente des troncs nerveux. Dans un petit nombre de réactions lépreuses, apparues sous forme d'érythème noueux, on a pu démontrer la présence de névrite infra-clinique.

Tous les moyens d'évaluation utilisés ont montré que la thalidomide était dotée d'un effet plus rapide que la prednisonne. A doses thérapeutiques faibles, la thalidomide semble également avoir été capable de prévenir les récurrences de réactions lépreuses. Les complications observées chez 5 malades n'ont pas pu être mises en relation avec la thalidomide; elles ont probablement été produites par la maladie sous-jacente dans 4 cas, et par un syndrome de Guillain-Barre dans le cinquième cas.

La signification, et les effets possibles, du MCV sont discutés.

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