Neuropathy of the somatic nervous system in rheumatoid arthritis is well documented. Peripheral neuropathy of sensory and sensori-motor types, multiple involvement of peripheral nerves (mononeuritis multiplex), digital neuropathy and lesions due to intrinsic pressure on single nerves are all described (e.g. Hart, Golding, and Mackenzie, 1957; Pallis and Scott, 1965). There is, however, a single report suggesting that interruption of the autonomic pathways may occur. Kalliomäki, Saarimaa, and Toivanen (1963) found that females with rheumatoid arthritis failed to sweat in response to an intradermal injection of nicotine on the forearms as compared with a matched group of control subjects. This failure appeared to be correlated with the presence of cold hands and a high erythrocyte sedimentation rate. No mention was made of clinical neuropathy in this study.

The present investigation was undertaken to determine the presence, extent, and location of autonomic nerve involvement in adult subjects with various types of rheumatoid neuropathy. Control subjects and patients with uncomplicated rheumatoid arthritis were also examined.

**Physiology of Sweating**

Sweating is an easily demonstrable autonomic function and is mediated by sympathetic nerves. The higher centres for thermo-regulatory sweating are generally thought to be situated in the hypothalamus (Hasama, 1930). The pathway of the long tracts is uncertain, but the preganglionic fibres, whose ganglion cells lie in the lateral horns between the first thoracic and first lumbar segments, emerge in the thoraco-lumbar outflow with the nerve roots and pass to the sympathetic ganglia where they terminate in a synapse. The cell body of the postganglionic neurone usually lies in one of the ganglia of the sympathetic chain. The postganglionic nerve emerges from the ganglion as the grey ramus communicans and passes to the periphery either with a main nerve trunk or in relation to a blood vessel (Fig. 1). In the dermis the nerve supplies sweat glands, erector pilae muscles, and the smaller blood vessels. The fibres divide terminally and a single nerve fibre may supply more than one structure and also act as a receptor for certain stimuli. Sympathetic nerves from T1—5 segments supply the upper limbs and from T5—L1 the lower limbs (Mitchell, 1953).

Postganglionic sympathetic fibres mediate the sympathetic axon reflex as described by Lewis and Marvin (1927). Antidromic impulses pass up the postganglionic fibre and produce a response in the effector organ. This is shown by the following experimental data:

1. Goose skin in response to faradic skin stimulation is abolished by extirpation of the stellate ganglion in cats (Lewis and Marvin, 1927).
2. Local sweating in response to faradic skin stimulation is abolished by a novocaine skin wheal at the site of stimulation, by injury to the brachial plexus, and by ganglionic sympathectomy, but remains intact after preganglionic sympathectomy. The response remains after
ANNALS OF THE REUHMATIC DISEASES

blocking a cutaneous nerve with local anaestheti
c at a point some distance proximal to its
termination. Hence the response is of the axon
reflex type and is mediated by postganglionic
sympathetic nerves (Bickford, 1938; Wilkins,
Newman, and Doupe, 1938).

(3) It has been shown in the cat that sweating and
piloerection cannot be elicited by electrical
stimulation of the distal portion of the severed
dorsal root, hence the response is not mediated
by sensory nerves. The response is, however,
obtained on stimulation of the ventral roots
(Collins and Weiner, 1961).

(4) It has been observed that intradermal injection
of acetyl choline or nicotine results in local
sweating and piloerection. This does not occur
after postganglionic sympathetic nerve degenera-
tion following ganglionectomy. It therefore
appears that sweating and piloerection which
follow an intradermal injection of nicotine or
acetyl choline are mediated by a sympathetic
axon reflex and that this response can be used to
test the integrity of postganglionic sympathetic
nerves in a manner analogous to that of histo-
mime and the sensory neurone (Rothman and
Coon, 1939; Coon and Rothman, 1940).

In normal subjects body heating causes symmetri-
cal sweating on the limbs and face (List and Peet,
1938), the magnitude of the response probably being
proportional to the density of sweat glands in the
area. Lesions of either the pre- or postganglionic
sympathetic pathways will result in deficient or
absent sweating. The integrity of the postgang-
lionic fibre can be tested by the cutaneous response
to an intradermal injection of nicotine or acetyl
choline, or to faradic stimulation. If the post-
ganglionic fibre supplying an area of deficient
thermo-regulatory sweating is intact, sweating and
piloerection will be seen, hence the thermo-regulatory
sweating abnormality must be due to involvement
of the preganglionic nerve. Absent sweating and
piloerection in the abnormal area signifies interrup-
tion of the postganglionic fibre, provided that
the presence of functioning sweat glands can be shown.
This can be demonstrated by the effect of local
heating to the area (Janowitz and Grossman, 1950).

Methods

The methods used were similar to those employed by
Bárány and Cooper (1956) in a study of diabetic patients.
Thermo-regulatory sweating was produced by placing
the subject in a bath of water at about 44° C. after the
limbs, which were supported above the water level, had
been covered with 5 per cent. castor oil in iodine paint
and sprayed with starch powder from an insufflato-
(Minor, 1927).

Subjects who were not already receiving salicylates in
large doses were given soluble aspirin 900 mg. one hour
before the test and all were given a hot cup of tea or
coffee immediately before the limbs were prepared. The
resulting sweat response appeared as blue-black dots
which later coalesced.

The areas in which sweating occurred after the subject
had been heated for 30 minutes were charted. The
maximum area of sweating was always apparent after
20 minutes' immersion. The only side effects of this
procedure were the appearance of a transient erythema-
tous rash on the limbs of three female patients in the
areas painted with iodine and a syncopal attack in one
female patient after getting out of the bath. During the
procedure the oral temperature rose by 1 to 3° F. and
tachycardia was frequently observed. None of the
subjects had severe anaemia or congestive heart failure.

The areas of skin in which sweating was either grossly
deficient or absent were then tested by faradic stimulation
and by intradermal injection of acetyl choline. Faradic
stimulation was provided by means of two silver elec-
rodes 3 mm. apart, placed on the skin after cleaning with
ether. The electrodes were connected to a standard
Multitone faradic stimulator, using a stimulus to pain
tolerance of as short a duration and of as great a
frequency as possible. In normal subjects (and in normal
areas in abnormal subjects) sweating was seen after
stimulation for 2 minutes. Its presence was detected by
applying quinizarin powder to the skin before stimulation.
Quinizarin, which turns deep purple where there is
sweating, was found to be more convenient than starch
and iodine for testing small areas of skin. Stimulation
in all test areas was continued for 4 minutes. A few
subjects failed to sweat in normal areas (i.e. where
thermo-regulatory sweating had been demonstrated)
probably because low pain tolerance prevented adequate
stimulation.

0·1 ml. 1 per cent. acetyl choline solution in normal
saline (preliminary experiments had shown this concen-
tration to give the maximal sweat response after 45 to
60 seconds) was injected intradermally into areas of
deficient sweating in all subjects. Nicotine was also used
in some subjects but was found to be less effective than
acetyl choline.

A sweat response to both faradism and to the intra-
dermal injection of acetyl choline was seen in an area
5 to 7 cm. in diameter around the point of stimulation.
The intensity of the response was greatest on the dorsum
of the hands, feet, and forearms, and was least on the
outer aspect of the legs. It appeared to be proportional
to the density of the sweat glands in the test areas.
In expressing results, the term "sweat loss" means completely
absent or grossly defective sweating.

In areas which showed both abnormal thermo-
regulatory sweating and absent sweating after faradism
and acetyl choline, the presence of sweat glands was
confirmed by heating the test area with a bottle containing
water at 80° C. for 2 minutes. This invariably produced
local sweating in all subjects.

In order to test the validity of these methods three
subjects with well documented neurological disorders
were examined:
AUTONOMIC NEUROPATHY IN RHEUMATOID ARTHRITIS

Sweating present in response to body heating. Absent sweating in response to body heating, acetylcholine, and faradism.

Fig. 2.—Sweating response. (a) Subject 1 (b) Subject 2

Subject 1. Syringomyelia with classical dissociated sensory loss on the arms and a right Horner’s syndrome (Fig. 2a).—Deficient sweat response occurred on the right arm and forearm and on the right side of the face. Local sweating occurred in these areas in response to faradic stimulation and intradermal acetyl choline injection. This was in accordance with the expected pattern and indicated damage to the preganglionic nerve.

Subject 2. Bilateral lumbar sympathectomy; right cervical (ganglionic) sympathectomy with surgically induced right Horner’s syndrome (Fig. 2b).—Sweating in response to faradic stimulation and intradermal acetyl choline injection did not occur in the abnormal areas, but did so in response to local heating. This indicated that the postganglionic nerve fibres were interrupted.

Subject 3. Vascular occlusion left middle finger; left cervical sympathectomy.—Thermo-regulatory sweating in this subject before sympathectomy showed a normal pattern and local tests showed no abnormality of the postganglionic fibres. 14 days after sympathectomy, thermo-regulatory sweating was almost completely abolished in the left arm, but the local sweat responses to acetyl choline and faradism remained normal; 6 months after sympathectomy these local responses had disappeared. This indicated that degeneration of the peripheral portion of the postganglionic nerve had to occur before the response disappeared. The local heating response remained normal.

Results

The numbers and types of patients examined by these methods are shown in Table I.

(1) Non-rheumatoid Controls

Eight male and five female subjects, who were healthy or who were suffering from diseases other

<table>
<thead>
<tr>
<th>TABLE I</th>
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<tbody>
<tr>
<td><strong>PATIENTS EXAMINED</strong></td>
</tr>
<tr>
<td>Diagnosis</td>
</tr>
<tr>
<td>(1) Healthy subjects or patients with diseases other than rheumatoid arthritis</td>
</tr>
<tr>
<td>(2) Rheumatoid arthritis complicated by neuropathy</td>
</tr>
<tr>
<td>(a) Symmetrical sensory neuropathy</td>
</tr>
<tr>
<td>(b) Lesions of major peripheral nerves</td>
</tr>
<tr>
<td>(3) Uncomplicated sero-positive rheumatoid arthritis</td>
</tr>
<tr>
<td>Total Patients Examined</td>
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</tbody>
</table>
than rheumatoid arthritis, were examined. They were of similar age to the patients with rheumatoid neuropathy.

In all except one the areas of deficient thermo-regulatory sweating were small and strictly symmetrical. In one subject, a female of 65 years with tophaceous gout, sweating failed to occur on the dorsum of the forearm and local tests with faradism and acetyl choline were negative. There was no sensory loss in the limbs of this patient.

(2) Patients with Rheumatoid Arthritis* complicated by Neuropathy

Two groups of patients with rheumatoid neuropathy have been examined, those with a symmetrical peripheral sensory neuropathy and those with a lesion of one or more main peripheral nerves. The

* All patients satisfied the American Rheumatism Association criteria for "definite rheumatoid arthritis" (Ropes, Bennett, Cobb, Jacox, and Jessar, 1959).

<table>
<thead>
<tr>
<th>Type of Neuropathy</th>
<th>Case No.</th>
<th>Sex</th>
<th>Sensory Loss</th>
<th>Pattern of Sweat Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Symmetrical Sensory</td>
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<td>M</td>
<td>Distal leg and knees, ankles and feet</td>
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</tr>
<tr>
<td>Neuropathy</td>
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<tr>
<td></td>
<td>4</td>
<td>M</td>
<td>Feet and ankles, recovering</td>
<td>Normal sweating in areas of sensory loss but postganglionic loss</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>M</td>
<td>Feet and ankles, recovered</td>
<td>Preganglionic type: lateral aspect one leg</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>F</td>
<td>Feet and hands</td>
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</tr>
<tr>
<td></td>
<td>7</td>
<td>F</td>
<td>Feet, recovering</td>
<td>Postganglionic: forearms only</td>
</tr>
<tr>
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<td>8</td>
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<td></td>
<td>11</td>
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<td></td>
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<td>Bilateral median</td>
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<td>15</td>
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<td>Posterior interosseus, recovered</td>
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AUTONOMIC NEUROPATHY IN RHEUMATOID ARTHRITIS

(i) Sweating Response

(ii) Sensory Loss

See Fig. 2 for key to sweating response

Fig. 3(a).—Case 6

Fig. 3(b).—Case 12

one or more fingers. Ten patients had at some time or other been noted as showing signs of digital sensory neuropathy (Pallis and Scott, 1965) and in five there was some digital sensory impairment at the time of sweat testing. In four of these the areas in which sweating was deficient corresponded to those in which there was sensory loss. In two patients there were areas of analgesia and sweat deficiency (of the postganglionic type) on the dorsum of the hand and these correspond with each other extremely closely. The significance of reduced sweating in the fingers of patients with rheumatoid neuropathy is difficult to assess, because areas of impaired sweating were demonstrated in the fingers of six of the thirteen control subjects, none of whom showed any sensory loss.
(3) Patients with Uncomplicated Sero-positive Rheumatoid Arthritis

One male and seven females with rheumatoid arthritis but with no clinical evidence of neuropathy were examined. Two of these patients showed abnormalities in sweat response outside the control range: both had patchy areas of sweat loss on the arms and legs (Fig. 4).

Fig. 4 (a and b).—Sweat loss in two patients with uncomplicated rheumatoid arthritis. See Fig. 2 for key.
**Discussion**

Although Kuno (1956) has emphasized the individual variation in sweating patterns which can occur in different subjects, or in the same subject under different conditions, our small number of non-rheumatoid controls did not show much variation. Of the eighteen patients with rheumatoid arthritis and neuropathy, fifteen showed abnormalities of sweating beyond the minor irregularities seen in control subjects. Those with the most extensive sensory neuropathy tended to show the greatest sweating disturbances. The area of sensory loss usually corresponded with that of sweat loss, though this was not always the case, and in three patients with recovering neuropathy (Cases 4, 5, and 7) the areas of sweat loss were small. There are at least two possible reasons why sensory loss does not always correspond to areas of deficient sweating. Firstly, there may be overlapping sympathetic innervation to the sweat glands. Guttmann (1940), studying sweat responses after complete division of single peripheral nerves, found that the resulting area of total sweat loss was much smaller than the area of sensory disturbance, and was surrounded by an intermediate zone with diminished response. Secondly, some sympathetic nerve fibres do not run with the major peripheral nerves but accompany blood vessels to the periphery. These fibres would escape lesions involving main peripheral nerves and would account for the small patches of sweating which were in fact always found on the finger tips and on some parts of the feet even in areas with complete analgesia. The converse may also be true, explaining the occasional disturbance of sweat response in the fingers and elsewhere in the absence of sensory disturbances.

In four patients (Cases 5, 8, 13, and 17) the pattern of sweat response was of preganglionic type, *i.e.* with deficient thermo-regulatory sweating but with acetyl choline and faradism responses present. But in all four the clinical neuropathy was recovering; and an alternative (and more likely) explanation to a preganglionic lesion is a postganglionic lesion with partial recovery, some response to acetyl choline and faradism returning before thermo-regulatory sweating can be demonstrated. We have not found any reports dealing with sweating responses during recovery and it appears that this should be investigated.

Postganglionic lesions were found in two subjects with severe sero-positive rheumatoid arthritis, neither of whom showed evidence of a sensory neuropathy though one had vascular lesions on the feet. A similar situation occurred in Case 18 where a pressure type of ulnar nerve lesion had recovered, but where extensive areas of sweat loss were present on the legs.

The significance of autonomic lesions in these three patients is uncertain. They possibly represent the onset of neuropathy which will later involve the sensory nerves; alternatively autonomic neuropathy may be a separate clinical entity with selective involvement of the non-medullated autonomic nerve fibres. Further studies on this type of patient are required, because autonomic neuropathy could account for some of the vasomotor disturbances seen in rheumatoid arthritis, as suggested by Kalliomäki and others (1963). Whether these patients will develop sensory neuropathy or extension of the autonomic neuropathy can be determined only by follow-up studies. It is possible that sweat tests may come to be of value in assessing the prognosis in certain patients.

All patients who showed evidence of autonomic neuropathy were well-established cases of rheumatoid arthritis, in many cases with evidence of arteritis. All except Case 18 had a positive sheep cell agglutination test. A series of patients with sero-negative rheumatoid arthritis has not, however, been examined, so that no conclusions can be drawn about the association of autonomic neuropathy with positive sheep cell tests.

The relation between clinical rheumatoid neuropathy and sero-positivity is established (Pallis and Scott, 1965), and, except where mononeuritis is due to local pressure from a swollen or damaged joint, the various syndromes of rheumatoid somatic peripheral neuropathy appear to be associated with, and probably caused by, vascular lesions and ischaemia. Our finding now of autonomic neuropathy in most of these patients suggests that there is a frequent and topographic association between the two types of neuropathy and that the mechanism of both is similar.

**Summary**

The thermo-regulatory sweating response to immersion in warm water and the local sweating response to intradermal injection of acetyl choline and faradic stimulation have been tested in eighteen patients with rheumatoid arthritis and peripheral neuropathy, in eight patients with uncomplicated rheumatoid arthritis, and in thirteen non-rheumatoid control subjects, in order to differentiate lesions of pre- and postganglionic fibres.

In the control group, areas of deficient sweating were small and symmetrical.

In most of the patients with rheumatoid arthritis and peripheral neuropathy, there was sweat loss in areas corresponding approximately to those of
cutaneous sensory impairment: this loss was usually of the type seen with interruption of postganglionic fibres, *i.e.* absent thermo-regulatory and local responses. In some patients the sweat response was normal. In four patients with thermo-regulatory sweat loss there was a positive local response suggestive of a preganglionic lesion, but in all four the clinical neuropathy was recovering or had recovered, and it is possible that the observed sweating abnormality represented a recovering postganglionic lesion rather than a preganglionic lesion.

Six of the eight patients with uncomplicated rheumatoid arthritis showed sweat responses similar to those of the control subjects, but two had larger areas of sweat loss: the significance of this is uncertain.

It is concluded that clinical sensory neuropathy in rheumatoid arthritis is usually accompanied by an autonomic neuropathy of postganglionic type corresponding to the sensory loss.

REFERENCES


Neuropathie autonome dans l'arthrite rhumatismales

RÉSUMÉ

Pour différencier entre les lésions des fibres pré- et post-ganglionnaires on étudia la réponse sudoripare thermo-régulatrice à l’immersion dans l’eau chaude et la réponse sudoripare locale à l’injection intradermique d’acétylcholine et à la stimulation paradigue chez 18 malades atteints d’arthrite rhumatismale et de neuropathie périphérique, chez 8 malades atteints d’arthrite rhumatismale incompliquée et chez 13 témoins non rhumatismaux.

Chez les témoins, les régions de suation insuffisante étaient petites et symétriques.

Dans la plupart des cas d’arthrite rhumatismale et de neuropathie périphérique, la sudation était abolie dans les régions correspondant à peu près à celles des alterations sensorielles cutanées; l’absence d’une réponse locale et thermo-régulatrice indiquait que la lésion était post-ganglionnaire. Chez quelques malades la réaction sudoripare était normale. Chez 4 malades la perte de la sudation thermo-régulatrice était accompagnée d’une réaction sudoripare locale positive, indiquant que la lésion était pré-ganglionnaire, mais chez tous les quatre, du point de vue clinique, la neuropathie était améliorée ou guérie; il est possible que l’anomalie sudoripare représente ici un lesions post-ganglionnaire en deye de guérison plutôt qu’une lésion pré-ganglionnaire.

Six malades sur huit, atteints d’arthrite rhumatismale incompliquée, ont accusé des réponses sudoripares similaires à celles des témoins, mais deux d’entre eux ont présenté de plus grandes surfaces de sudation; on ne sait pas exactement ce que cela signifie.

On conclut que la neuropathie sensorielle clinique dans l’arthrite rhumatismale est généralement accompagnée d’une neuropathie autonome du type post-ganglionnaire à territoire commun.

Neuropatía autónoma en la artritis reumatoide

SUMARIO

Para diferenciar entre lesiones de fibras pre y postganglionares se estudió la respuesta sudoripara termo-reguladora a la inmersión en el agua caliente y la respuesta sudoripara local a la inyección intradérmica de acetilcolina y a la estimulación faradica en 18 enfermos con artritis reumatoide y neuropatía periférica, en 8 enfermos con artritis reumatoide sin complicación y en 13 testigos sin reumatismo.

En los testigos, regiones de exudación deficiente fueron pequeñas y simétricas.

En la mayoría de los casos de artritis reumatoide y de neuropatía periférica la exudación fue perdida en regiones que correspondieron aproximadamente a las con alteraciones sensoriales cutáneas; la reacción sudoripara local y termo-reguladora negativa indicó que la lesión fue post-ganglionar. En algunos enfermos la reacción sudoripara fue normal. En 4 enfermos la pérdida de la exudación termo-reguladora se vio acompañada de una reacción sudoripara local positiva sugeriendo una lesión pre-ganglionar, pero en todos ellos, clínicamente, se observó el restablecimiento parcial o completo de la neuropatía, de modo que se puede tratar aquí de mejorías de lesiones post-ganglionares.

De ocho enfermos con artritis reumatoide sin complicaciones, seis acusaron reacciones sudoriparas similares a las de los testigos, pero dos otros presentaron mayores zonas de exudación; no es cierto el significado de esto.

Se concluye que la neuropatía sensorial clínica en la artritis reumatoide se acompaña generalmente de neuropatía autónoma del tipo post-ganglionar de distribución común.