ARSENICAL ENCEPHALOPATHY IN INDIAN TROOPS

BY

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Introduction

Untoward effects of antisyphilitic treatment, described as "arsenical encephalopathy," have been observed ever since trivalent arsenical compounds have been in use for the treatment of syphilis. Paul Ehrlich as early as 1914 believed the essential cause of the condition to be a dilatation of the small blood vessels of the brain which is followed by edema and perivascular bleeding. He believes that the vascular change was due to some derivative of arsphenamine formed inside the body, as is suggested by the delay of several days before the characteristic reaction took place. Non-hæmorrhagic cases with necrosis and softening were described by Pollack and Riehl (1930). Demyelinating lesions were found by Dorothy Russell (1937). A comprehensive survey of the literature was made by Glaser and Immerman (1935). The condition has also occurred when the trivalent arsenicals have been used in conditions other than syphilis.

During 1943 and 1944 there were many fatalities among Indian troops undergoing treatment of syphilis with arsenicals. At the Third Medical Division's Conference, Southern Army, Poona, 1944, Krainer reported seventeen cases under the heading "arsenical encephalopathy."

In view of this high incidence an attempt was made to outline and classify the clinical picture and pathology of this condition with special reference to factors responsible for its appearance among Indian personnel. Reports of 184 cases were received from all over India. These included 37 cases observed in the Indian Military Hospital, Dunkirk.

Analysis of Material

An analysis of available case histories led to the following conclusions:

1. The incidence of arsenical encephalopathy in Indian troops is higher than the incidence rate of classical hæmorrhagic encephalopathy.

2. The incidence of this condition in Indian troops treated with weekly injections of N.A.B. is of the same order as the incidence in Indian civilians.

3. The increased incidence rate of this condition observed since 1943 is essentially the effect of treatment with bi-weekly, as opposed to the old standard treatment with weekly injections. (The dosage of arsenicals in the two treatment groups was uniformly standardized. Patients weighing 115 lb. and above received alternating 0.3 and 0.45 g. of N.A.B. Patients below 115 lb. received 0.3 g. per injection.)

4. The increased incidence after bi-weekly injections refers to Madrasiss and Mahrattas only, the incidence in other provincial groups showing no significant increase. It is noteworthy that the incidence rate is higher in the two provincial groups with the lower average body weight.

5. Malaria during or immediately before antisyphilitic treatment is an important factor in increasing the rate of arsenical encephalopathy.

Statistics.—The reported incidence of arsenical encephalopathy in the Army in India from August, 1943, to March, 1945, was:

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>Indian cases</td>
<td>181</td>
</tr>
<tr>
<td>British cases</td>
<td>3</td>
</tr>
</tbody>
</table>

The Indian cases were distributed as follows:

- Madrassiss: 97
- Mahrattas: 22
- Others: 62

Total: 181

Table I shows the total number of cases undergoing antisyphilitic treatment, and the incidence of arsenical encephalopathy, in the Indian Military Hospital, Dunkirk, and the Indian Military Hospital, Jalahall. The average incidence rate in these groups is far higher than that recorded in the literature, for example, 0.2 per thousand in Glaser and Immerman's statistics. Statistics from the Madras General Hospital (1,000 cases treated with neoarlesphenamines: Rajam and Rao, 1939) show a rate of 0.5 per cent., that is, twenty-five times Glaser and Immerman's figure.

Table II shows the incidence of arsenical encephalopathy classified according to weekly and bi-weekly injections. It shows that the incidence rate of arsenical encephalopathy is not due to a proportional increase in all groups. The incidence is six times the incidence following weekly injections in Madrasiss and four times in Mahrattas, but it is only 0.2 per cent. higher in Indians from other provinces, which is below statistical significance.
TABLE I

<table>
<thead>
<tr>
<th>Provincial group</th>
<th>Syphilis under AST</th>
<th>Cases of arsenical encephalopathy</th>
<th>Incidence %</th>
</tr>
</thead>
<tbody>
<tr>
<td>I.M.H. Dunkirk</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Madrassis</td>
<td>657</td>
<td>14</td>
<td>2·1</td>
</tr>
<tr>
<td>Maharattas</td>
<td>460</td>
<td>16</td>
<td>3·5</td>
</tr>
<tr>
<td>Others</td>
<td>792</td>
<td>7</td>
<td>0·9</td>
</tr>
<tr>
<td>Total</td>
<td>1,909</td>
<td>37</td>
<td>1·9</td>
</tr>
<tr>
<td>I.M.H. Jalalahalli</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Madrassis</td>
<td>1,548</td>
<td>31</td>
<td>2·0</td>
</tr>
<tr>
<td>Maharattas (not separately recorded)</td>
<td>820</td>
<td>8</td>
<td>1·0</td>
</tr>
<tr>
<td>Total</td>
<td>2,368</td>
<td>39</td>
<td>1·6</td>
</tr>
</tbody>
</table>

Comparison of figures for the Indian Military Hospital, Dunkirk, and the Indian Military Hospital, Jalalahalli, from January, 1944, to March, 1945.

The average weight of Madrassis and Maharattas is below that of the group classified as "Others" (see Table III).

Social Incidence.—Of the patients under antisyphilitic treatment, 11·1 per cent. (212 out of 1,909 cases) were non-commissioned officers; only three of these developed arsenical encephalopathy. No case was recorded in a Viceroy-commissioned officer or gazetted Indian officer. It was considered that at least part of the high racial incidence in Indians is social incidence. It seems a reasonable conjecture that a deficient diet plays a part. Lydon (1944) has stressed the possible relationship of Vitamin B1 and C deficiency. The poor Madrassi diet is known for its deficiencies (McCarrison, 1921).

Malaria.—Thirteen cases out of a series of 1,909 developed benign tertian malaria during or shortly before undergoing antisyphilitic treatment; of these, four developed arsenical encephalopathy, giving an incidence of 30·8 per cent. as against 1·9 per cent.

Time of Onset of Arsenical Encephalopathy.—The intervals between the first injection of N.A.B. and the onset of arsenical encephalopathy can be seen in graphs J, 2, 3, and 4.

The influence of the stage of the syphilitic infection on the incidence rate of arsenical encephalopathy is shown in Table IV, from which it can be seen that the incidence of arsenical encephalopathy in sero-positive primary and secondary syphilis is no higher than in sero-negative primary syphilis. On the contrary, the incidence rate in sero-negative primary syphilis is higher than in the other two groups of this series, the difference being below statistical significance.

Mortality Rate.—The mortality rate is shown in Table V. The mortality rate of arsenical encephalopathy was 76 per cent. in the series of Glaser and Imerman; it will be seen, therefore, that the mortality rate is lower in Indian troops, but the fatalities from encephalopathy in Indian troops are 75·9 per cent. of all fatalities during antisyphilitic treatment, as against 50 per cent. in the series of Cole and others (1931).

The Clinical Picture

This description is based upon the analysis of 37 cases observed in the Indian Military Hospital, Dunkirk. In general, a fairly definite sequence of events could be outlined.

Prodromal Symptoms.—In a few cases, several days before the onset the patient exhibited headache, fever with a negative blood slide, and malaise. This was often associated with excessive thirst, a feeling of chilliness and exhaustion, and, on occasion, abdominal colic and epigastric tenderness. These became enough to warn us that encephalopathy threatened.

Onset.—Gradual and sudden types occurred in approximately equal numbers; the latter commenced as an epileptic fit followed by evidence of cerebral irritation.

Progress.—For the purpose of description four subsequent stages may be distinguished. For the sake of convenience they may be called, somewhat arbitrarily, stage one, two, three, and coma. A patient may never progress beyond stage one or stage two.

The essential feature of stage one is that the patient, though mentally retarded, is still accessible and is able to walk about. He may complain of malaise and lethargy, and often of fear; visual hallucinations and confusion may be prominent. He tends to wander from his bed. On examination the pupils are found to be

TABLE II

<table>
<thead>
<tr>
<th>Frequency of injections</th>
<th>No. of cases of AST</th>
<th>Number of A.E.</th>
<th>Incidence rate %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Madrassis</td>
<td>Maharattas</td>
<td>Others</td>
</tr>
<tr>
<td>Weekly</td>
<td>342</td>
<td>173</td>
<td>380</td>
</tr>
<tr>
<td>Bi-weekly</td>
<td>315</td>
<td>287</td>
<td>412</td>
</tr>
</tbody>
</table>

Incidence of arsenical encephalopathy classified according to frequency of injections.
Classification of cases under antisyphilitic treatment and cases of arsenical encephalopathy, according to provincial group and body weight.

small and to react to light. There may be a general hyperesthesia, but plantar reflexes and deep reflexes are normal while the abdominals may be absent. Questions are answered correctly, but there is a noticeable delay.

In the second stage the patient is more drowsy but can be roused. He is, however, unco-operative and resists examination. There is often hyperesthesia and fever. Examination reveals small irregular pupils reacting to light. The deep reflexes are increased with an upgoing toe. The superficial reflexes are lost and there is an upgoing toe.

The third stage is always pyrexial with increased respiratory rate, stertor, hiccough, and loss of sphincter control. There are frequent fits resembling status epilepticus. There may be teeth-grinding, moaning, etc. Examination shows dilated pupils reacting sluggish to light, later becoming fixed. The deep reflexes and the swallowing reflex are still present. Kernig’s and Babinski’s sign may be present.

This is followed by deep coma in which the respiratory rate may exceed sixty per minute. The deep reflexes are lost. The pupils are fixed, and there is a tendency to hyperpyrexia. Every case which reached this stage died.

Special Examinations.—Examination of the fundus in ten cases showed no abnormalities, and blood pressure was within normal limits, the average being 117/73 mm. Hg. In ten cases that had reached stage three the temperature was 98-4° F., the pulse rate 84, and the respiration rate 26 per minute. Babinski’s sign was negative. By Oct. 14 he was dazed, refused to speak, and made occasional twitching movements with the

The influence of the stage of the syphilitic infection on the incidence rate of arsenical encephalopathy.

cerebrospinal fluid pressure varied from 45 to 190 mm. water, the average being 87. The cerebrospinal fluid cell count ranged from 0 to 40, being under 4 in 32 cases, from 5 to 10 in 3 cases, and 11 and above in 2 cases. The total protein ranged from 70 to 900 mg. per 100 c.cm., average 207 mg. It was noted that no case with over 300 mg. per 100 c.cm. survived.

The cerebrospinal fluid Wassermann reaction was negative in 17 cases and positive in 3.

The white cell count ranged from 5,000 to 21,000 per c.mm.

Albuminuria was present in 12 out of 37 cases.

Treatment

Treatment consisted of general nursing and a standard treatment as recommended by Army Directive, which included: venesection, lumbar puncture, adrenaline, morphia, atropine, and the routine use of intravenous calcium gluconate. The results of this treatment as well as of additional procedures—administration of sodium thiosulphate and oxygen, were inconclusive. Case records illustrative of the types of cases encountered are appended.

Case Reports

Case 1.—A Madrassi Havildar (Sergeant), aged 19 years, weight 90 lb., was admitted to hospital on Sept. 17, 1944, with a generalized papular rash of 8 days' duration and a sore in the right inguinal region. Blood Wassermann and Kahn reactions were negative. He received four injections of N.A.B. bi-weekly, to a total of 1-2 g.

On Oct. 13, eleven days after the first injection he was apprehensive, restless, and groaning, but conscious and able to answer questions. He was incontinent of urine. The temperature was 98-4° F., the pulse rate 84, and the respiration rate 26 per minute. Babinski’s sign was negative. By Oct. 14 he was dazed, refused to speak, and made occasional twitching movements with the
fingers. The left pupil was larger than the right, both reacting to light; there was bilateral nystagmus. The deep reflexes were present, and the plantar reflexes extensor. He was still incontinent of urine. The cerebrospinal fluid protein was 100 mg. per 100 c.cm.

By Oct. 16 the patient had recovered. This was a mild case, with fear as an early symptom.

Case 2.—A Madrassi Sepoy, aged 22 years, weight 95 lb., who had had a penile sore one year previously and now had a scar from the old chancre, and whose blood Wassermann and Kahn reactions were positive, was given four injections of N.A.B. bi-weekly, totalling 1-2 g. On Sept. 12, 1944, twelve days after the first injection, he suddenly fell while on fatigue. He was unconscious, and had hematemesis. The temperature was 100° F., and he frothed at the mouth. There were choroidal movements of limbs and head, and athetosis of the hands. The eyes showed conjugate deviation to the left. The pupils were moderately dilated and equal, and reacted to light. Babinski’s reflex was negative. The deep reflexes were sluggish, and the abdominal reflexes absent. The right plantar reflex was extensor, the left equivocal. There was incontinence of urine. The cerebrospinal fluid pressure was 70 mm. water, five cells, protein 100 mg. per 100 c.cm.

On Sept. 13 the patient was grinding his teeth and had occasional convulsions. On Sept. 15 he was conscious and responded to questions, but with some delay. Abdominal reflexes were absent and plantar reflexes flexor; he was incontinent of urine on this day and the next, but by Sept. 19 he felt well.

This was a typical case with sudden onset.

Case 3.—A Sepoy, aged 36, weight 120 lb., classified as “Others,” was admitted to hospital on June 4, 1944, with inguinal bubo. Routine blood Wassermann and Kahn examinations were positive. He received five injections of N.A.B. at weekly intervals, total 1-8 g.

On July 6, seventeen days after the first injection, he was found wandering aimlessly and was conducted back to the ward. The next day he looked vacant and was unable to reply to questions; five hours later he was found unconscious. The temperature was 102° F., the respiration rate 36 per minute, and he was bathed in sweat. Examination of the eyes showed the pupils moderately dilated, reacting to light, left external rectus palsy, and normal fundi. Babinski’s sign was positive. Jaw clonus was present, and there was fibrillation of masseters and temporales, but no paresis. There were intermittent clonic movements of the right arm and of both legs, with continuous athetoid movements of hands and feet. The deep reflexes were increased, the abdominal reflexes absent, and the plantar reflexes extensor. There was incontinence of urine. The blood pressure was 140/80 mm. Hg, and the cerebrospinal fluid contained 210 mg. of protein per 100 c.cm.

By July 8 the patient was in deep coma, with pupils contracted and fixed, and all deep reflexes absent. His condition rapidly worsened, and he died.

This case demonstrates the “three stages” preceding the coma which leads to death.

Case 4.—A Pioneer, age and weight unknown, classified as “Others,” was admitted on April 4, 1944, to the venereal disease centre, Serampore. The blood Wassermann and Kahn reactions were positive. He received three injections of “Evarsan” at bi-weekly intervals, a total of 9-9 g., and had no reaction.

On April 18 the blood Kahn reaction was still positive. On April 25, thirteen days after the first injection, the patient took a normal meal and then collapsed where he was sitting, after which he was unable to answer questions. He was not unconscious, but stupid (stage two). He received 1 c.c.m. of adrenaline intramuscularly at 14.40 hours, the dose being repeated at 15.10 hours when he also had 5 per cent. glucose saline, 20 c.c.m., and calcium gluconate, 10 c.c.m. intramuscularly, and a lumbar puncture. He improved, and next day was fully conscious but weak; he was given glucose and milk and kept in bed.

On April 27 he was found collapsed, bathed in sweat, and quite unconscious at approximately 06.15 hours. He was given 1 c.c.m. of adrenaline intramuscularly at 06.30 hours, at 08.30 hours, and thereafter at three-hourly intervals; also 50 per cent. glucose, 20 c.c.m., intravenously at 07.00 hours, and 100 c.c.m. at 15.00 hours.

### Table V

<table>
<thead>
<tr>
<th>Complications of AST</th>
<th>No. of cases</th>
<th>Fatal cases</th>
<th>Fatality rate %</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.E. ....</td>
<td>37</td>
<td>20</td>
<td>54.1</td>
</tr>
<tr>
<td>Other than A.E. ...</td>
<td>49</td>
<td>6</td>
<td>12.2</td>
</tr>
</tbody>
</table>

**Mortality**

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>No. of cases</th>
<th>Interval between last injection and death in days</th>
<th>Total N.A.B. given in g.</th>
<th>Arsenic contents in mg. %</th>
<th>Arsenic content of brain in % of liver</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arsenical encephalopathy</td>
<td>6</td>
<td>1-4</td>
<td>1.35</td>
<td>0.17</td>
<td>0.58</td>
</tr>
<tr>
<td>Other fatalities during AST</td>
<td>3</td>
<td>1-5</td>
<td>2.95</td>
<td>0.026</td>
<td>0.3</td>
</tr>
</tbody>
</table>

Arsenic contents of tissues; average values.
A diagnosis of cirrhosis was made. Pseudolobules and there was fibrous tissue in the round-cell infiltration. \( \text{Reticulum stain showed} \) cells, arteries, and adhesions were detected. \( \text{Histological examination of the liver showed focal round-cell infiltration with} \) peripheral histiocytes and fibroblastic proliferation and occasional central necrosis; a few histiocytic giant cells, and also symplasmic giant cells formed by degenerating liver cells, were seen. \( \text{Reticulin stain showed} \) overgrowth of delicate reticulum fibres in the areas of necrosis. There was regeneration of liver cells in the neighbourhood under formation of pseudolobules and there were also foci of scar formation. A diagnosis of syphilitic hepatitis with early syphilitic cirrhosis was made.

The medulla oblongata showed occasional perivascular round cells, but no other pathological changes were detected.

In the midbrain there was mild thickening of the meninges, mild proliferation of the astroglia subpial, and perivascular necrosis around some of the large rami perforantes, with demyelination. In the cerebellum there was hyperemic congestion of the meninges. In the cortex the meninges were moderately thickened, with scanty round-cell infiltration. The meningeal vessels were congested, and there was hyperemic congestion of the cortex and of the subcortical white matter. There was no evidence of hemorrhage. The round cells in the meninges were lymphocytes and histiocytes. This was a case of relapse.

Case 5.—A Sepoy, age and weight unknown, was admitted to the venereal disease centre on Dec. 9, 1943, with a penile sore. The blood Wassermann was strongly positive, the Kahn positive. He had two injections of N.A.B. at a week's interval, total 0.6 g.

On Dec. 26, eleven days after the first injection, he was found to be behaving abnormally. Pupils were normal and reacted to light. There was retention of urine but no rise of temperature. He died eight hours later.

 Necropsy showed an Indian male of average build, with bloodstained froth round the mouth. The pleura was free, but there was œdema of the lungs. Heart, abdominal cavity, liver, spleen, bladder, and prostate were normal. The brain showed confluent punctate hemorrhages and hemorrhagic softening in the pes of the pons.

Histological examination of the cortex and subcortical
white matter showed hyperæmic congestion. The subcortical vessels showed severe distention of the perivascular space by homogeneous viscous proteinic material. There were scanty round cells perivascularly. Edema was present. The pons pontis showed engorgement of the large vessels and changes of the same type as described in the cortex. There was also extensive confluent capillary haemorrhage of the ring type, and occasional perivascular haemorrhage. Extensive necrotic changes existed in the neighbouring brain substance with a tendency towards homogenization. There was edema. The liver showed hyperæmic congestion, occasional fatty change, and marked periportal round-cell infiltration. The spleen showed haemorrhagic patches in the pulp. This was a case of "haemorrhagic arsenical encephalopathy."

Pathology

This report is based on nineteen post-mortem examinations with subsequent histological examination. The central nervous system and internal organs of forty-one additional cases were also examined histologically. In view of the fact that the histological lesions are the same as in classical "haemorrhagic arsenical encephalopathy" and differ only in numbers and intensity, the account will be brief.

Macroscopic Examination

External Appearance.—A healing or healed penile sore was a common feature. Skull and Brain.—Congestion of the meningeal veins was a constant feature. The most frequent findings were: a mild swelling of the brain, a pale appearance of the white matter; the grey matter was either normal or congested. The brain was not "wet." Only one case showed a pressure cone (of the cerebellum), and only two showed the appearance of "pink brain." Subarachnoid ecchymoses were seen occasionally. Chest.—There was usually hypostatic congestion of the lungs.

Other Organs.—The spleen was commonly enlarged; malarial pigmentation was found only in one case, while smears yielded negative results for malarial parasites.

Histological Examination

(The number of cases examined is given in parentheses). Liver (45).—Fatty changes were a common feature; small haemorrhages, focal fatty degeneration, and foci of liver cell necrobiosis were seen occasionally. Specific gummatus lesions were seen in three cases only (Fig. 1). Periportal infiltration with lymphocytes, plasma cells, and occasional eosinophils was a prominent and nearly constant feature. Spleen (30).—Cases with splenic enlargement showed congestion, haemorrhage, and a moderate degree of pulp hyperplasia. Suprarenals (20).—Fatty infiltration of the cortex was commonly found. Kidneys (26).—The commonest findings were hyperæmia, congestion, and cloudy swelling. Central Nervous System (60).—The entire nervous system was examined systematically. There was only a small number of histological lesions, and they were usually of low-grade intensity. The brain stem and white matter were as a rule more involved than the cortex and the medulla oblongata. Congestion and stasis were a constant feature—predominantly involving the veins and the arterioles. Haemorrhages were scanty and usually only perivascular. Capillary haemorrhage of the "ring" type was found only rarely (Fig. 2). There was normally no direct evidence of endothelial damage (Fig. 3). Thrombotic changes were rare (Fig. 4). A viscous exudation of plasma, perivenuous and perilarietal (but not around capillaries) was a fairly constant finding (Fig. 5).

Inflammatory changes in the form of perivascular round-cell infiltration were seen in nearly all cases (Fig. 6). Patchy round-cell infiltration of the meninges was also seen (Figs. 7 and 8). Unlike the vascular changes, the round-cell infiltration was also commonly found in the cortex. All the inflammatory changes were mild. The inflammatory cells were lymphocytes, histiocytes, and plasma cells. Parenchymatous changes were also mild (Fig. 9), and were partly secondary to perivascular haemorrhage and viscous exudation. Nerve-cell changes were absent. Though all the different types of lesions followed the same pattern of distribution, a considerable independence of individual lesions was observed; perivascular and round-cell infiltration were seen without haemorrhage, and viscous exudation independent of haemorrhage. There was mild interstitial oedema. (Gross oedema with its macroscopical signs was absent.) Some cases showed atypical features. Two showed haemorrhagic features—one of them in the form of purpuric capillary haemorrhages of the "ring" type in the subcortical white matter, and the other in the form...
of a confluent capillary and perivascular hemorrhage in the pons. Two cases showed definite perivascular demyelinating lesions not secondary to viscous exudation or hemorrhage. The lesions were situated in the occipital white matter in one case, and in the midbrain in the other (Fig. 10).

**ARSENIC CONTENTS OF TISSUES.**—These were determined on tissue digests with the Gutzeit method. Table VI shows the average values obtained. Our values are in good agreement with those recorded by Osterberg and Kernohan (1934).

**Comparison of Arsenical Encephalopathy in Indian Troops with Classical Hemorrhagic Encephalopathy**

Comparison of "arsenical encephalopathy in Indian troops," as described here, with typical "hemorrhagic encephalopathy," as recorded in the literature, shows a seemingly adequate pathological basis in the latter group, and macroscopical and histological findings insufficient to explain the severity of the clinical picture in the former. The pathological relationship between the two conditions is stressed by the occasional occurrence of typical hemorrhagic cases in this series, and by two cases with typical demyelinating lesions. It is noteworthy that fully developed demyelinating lesions were seen by Dorothy Russell only in cases of four and six days' duration. One of our cases presenting fully developed demyelination is a case of "relapse" of 71 hours' duration from the onset to the fatal outcome; the second case was of 36 hours' duration only. (The interpretation of this is somewhat confused; as the patient had been vaccinated four days before the onset of arsenical encephalopathy; his reaction to vaccination was "modified positive.")

The possibility that demyelinating lesions observed in cases of considerable duration—4 to 6 days in the series of Dorothy Russell, 71 hours and 36 hours in this series—are the histological manifestation of functional alterations at corresponding sites, that is, the perivenous and periarterial white matter, in cases of shorter duration, has been considered. Occasional perivascular swelling of myelin sheaths and axis cylinders, independent from hemorrhage, as well as an occasional minute focus of early perivascular necrosis, is recorded by Dorothy Russell (1937) in one case of classical hemorrhagic arsenical encephalopathy of six hours' duration.

Arsenical encephalopathy in Indian troops is the "pathological equivalent" of classical "hemorrhagic encephalopathy."

Non-hemorrhagic cases of arsenical encephalopathy are recorded occasionally in the literature. It seems that the number of typical hemorrhagic cases under record is somewhat exaggerated, as there is a tendency to regard cases with a few perivascular hemorrhages as examples of it. In Halcrow's (1943) case "the lesions (hemorrhages and others) present were not numerous, therefore careful searching was necessary. Some blocks of brain tissue revealed no lesions."

Visible damage to the capillary endothelium with subsequent thrombosis and rhesis of blood vessels is far more prominent in hemorrhagic arsenical encephalopathy than in arsenical encephalopathy observed in Indian troops. Viscous perivascular exudation was seen perivenously and periarterially. It indicates loss of selective permeability of blood vessels. It is of interest that viscous perivascular exudation was seen in the absence of gross edema. In cases of arsenical encephalopathy with insufficient pathological findings recorded in the literature, gross edema was regarded as the cause of the clinical picture and fatal outcome. The examination of this series shows that gross edema sufficient to produce increased intracranial pressure is not a feature of the condition.

The macroscopical and histological findings of arsenical encephalopathy in Indian troops are interpreted here rather as signs than as the cause of the clinical picture.

The presence of a circulatory disturbance is indicated by stasis, congestion, and perivascular and capillary hemorrhage; viscous perivascular exudation indicating the loss of selective permeability of blood vessels.

Comparison of the pathological findings and of the clinical picture in this series with the findings in classical "hemorrhagic encephalopathy" raises some doubt that hemorrhage itself is the only cause of the clinical picture in hemorrhagic encephalopathy, as the clinical picture may develop in the absence of gross or purpuric hemorrhage.

Inflammatory changes are more constant in this series than in classical hemorrhagic encephalopathy, but the inflammatory changes are only mild. Koenigstein and Spiegel (1920) have referred to the difficulty of distinguishing between inflammatory lesions due to syphilis and inflammatory lesions due to arsenical encephalopathy in hemorrhagic cases. The evidence of mild thickening of blood vessels with obvious perivascular round-cell infiltration, the coexistence of patchy meningeal round-cell infiltration, and the predominantly histiocytic character of the inflammatory cells, suggest in our opinion that at least part of the inflammatory lesions is the manifestation of preclinical neurosyphilis. In good agreement with this interpretation of the inflammatory lesions is the evidence in the literature that a large proportion of the cases of syphilis show abnormalities in the fluid early in the disease " (Merritt and Fremont-Smith, 1938). It is pointed
Fig. 1.—Liver; small gummatus periportal lesions.

Fig. 2.—Pons; capillary haemorrhages of the ring type.

Fig. 3.—Cerebellum; meningeal ecchymosis, capillary vasodilatation with swelling of endothelial cells, and capillary haemorrhages.

Fig. 4.—Pons; incomplete thrombosis of a venule.

Fig. 5.—Subcortical white matter from the same case as in Fig. 4; perivascular viscous exudation.
Fig. 6—Pons: perivascular round-cell infiltration.

Fig. 7—Cortex: meningeal round-cell infiltration.

Fig. 8—Enlargement of Fig. 7: lymphocytes and histiocytes.

Fig. 9—Subcortical white matter: perivascular neurogliaproliferation; presence of mobile cells.

Fig. 10—Midbrain: perivascular necrosis and denervation.
out that none of our cases shows gross evidence of meningo-vascular syphilis.

Specific lesions in the internal organs other than the central nervous system were confined to the liver. Three cases show gumma formation, and one case syphilitic hepatitis. Marked periportal round-cell infiltration was interpreted as a manifestation of syphilis.

**Aetiology**

It seems that the facts known and some additional facts emerging from this investigation are insufficient to form the basis of a satisfactory and comprehensive explanation as to the aetiology of arsenical encephalopathy. For this reason only a short discussion of the various theories in being, and interpolation of the findings recorded in this investigation, is attempted here, the guiding principle of the discussion being to separate so far as possible the cause of the condition from its mechanism.

**Cause**

Activation of a pre-existing virus has been considered as the cause of arsenical encephalopathy. In other words, arsenical encephalitis would be a virus disease, the virus being pre-existent in the body of the individual and activated by the administration of a trivalent organic arsenical compound. This hypothesis bases its claims on the similarity of the picture of acute perivasculary myelinolysis and haemorrhagic encephalitis following various virus infections or antirabic treatment, with the picture of haemorrhagic and demyelinating arsenical encephalopathy. It is admitted that the exact relationship between the original virus infections and the subsequent encephalitis in the group of diseases referred to is not known. No additional facts have been found here to give more weight to the infection theory, so far as activation of a pre-existing virus is concerned.

Sterilization of syringes after each injection had no influence on the incidence rate of arsenical encephalopathy, and when two cases of arsenical encephalopathy occurred on the same day a subsequent check on the order in which the patients were injected revealed that the patients belonged to different batches. There seems to be no proof that a virus is carried by the syringe.

Various authors have pointed to the complex of Jarisch Herxheimer's reaction as the cause of arsenical encephalopathy. Our material gives no evidence in this direction. If the condition is Jarisch Herxheimer's reaction it should follow twenty-four hours after the first injection of the N.A.B., and if this be dispensed with one would expect that the cerebrospinal fluid would be Wassermann-positive in these cases at the time of the arsenical encephalopathy. Out of twenty-five cases examined, the cerebrospinal fluid Wassermann was negative in 17 and positive in 8. For that reason it is thought that Jarisch Herxheimer's reaction is not the cause of arsenical encephalopathy in this series.

Authors who believe that acute perivascular myelinolysis following various infections is the effect of sensitization, and not a virus disease in itself, expressed the opinion that arsenical encephalopathy is the result of sensitization against arsenic. Various points are in favour of this assumption, as the time of onset of arsenical encephalopathy after the first injection is the same as the common time of onset of acute perivascular myelinolysis or haemorrhagic encephalitis after the original virus infection; also the interval between the onset of serum sickness and administration of serum is of comparable order. Trivalent organic arsenical compounds, especially if administered together with serum, produced sensitization and anaphylactic reaction in guinea-pigs (Landsteiner and Jacobs, 1936). For that reason sensitization cannot be excluded beyond doubt as a causative factor of arsenical encephalopathy. Reports of untoward accidents following administration of arsenic compounds in patients who survived an attack of arsenical encephalopathy are recorded in the literature. An observation to the contrary was put at our disposal by Captain L. Cohen, R.A.M.C. After having had an attack of arsenical encephalopathy a patient received erroneously an injection of N.A.B., with no ill effects. A second case treated with intramuscular injections of a trivalent arsenical compound subsequent to an attack of arsenical encephalopathy showed also no untoward effects.

Finally, toxic effects of trivalent organic arsenical compounds have also been regarded as the cause of arsenical encephalopathy. Osterberg and Kernohan (1934) reported "relatively high arsenic contents," especially of the white matter of the cerebrum, in cases of haemorrhagic arsenical encephalopathy. "It has been suggested by Voegtlin and his co-workers that the same mechanism by which the arsenicals are lethal to parasites (spirochaetes) is apparently concerned in their toxicity for the tissues of the host" (quoted from Goodman and Gilman, 1941).

After one injection of arsphenamine 50 to 70 per cent. of the arsenic is excreted within one week, and this is a determinant of the weekly interval between injections in patients (Goodman and Gilman, 1941). The standard treatment of syphilis introduced on this basis—weekly injections of trivalent organic arsenicals—gives a high percentage of cures with a minimum of untoward effects and fatalities.
ARSENICAL ENCEPHALOPATHY IN INDIAN TROOPS

In the endeavour to shorten the time of treatment and to ensure a higher percentage of cures of syphilis, massive and continuous treatment with N.A.B. and mapharside was introduced. Increased spiroidal toxicity obtained by these means was accompanied by increased tissue toxicity.

The Army treatment of syphilis with bi-weekly injections is a compromise between massive treatment and standard treatment with weekly injections. In this investigation it was found that the incidence rate of arsenical encephalopathy in certain groups of Indians—Madrasis and Mahrattas—was significantly higher with bi-weekly than with weekly injections; this fact is relevant to any hypothesis of causation.

It is often recorded in the literature that arsenical encephalopathy is as common with high as with low doses of N.A.B. Our material does not allow any statement to be made to this effect, as the dosage was standardized according to body weight and for that reason comparable in the different dosage groups.

The following line of thought has been taken here. The basis of the weekly standard treatment was the knowledge that a high percentage of arsenic (50 to 70 per cent.) is excreted within one week after injection. Treatment with weekly injections avoids accumulation of arsenic. It appears logical that if treatment is given at bi-weekly intervals, with either the same dose or even with a somewhat lower dose of N.A.B., a higher level of arsenic will be maintained for a longer time. As bi-weekly treatment is apparently associated with an increased incidence rate of arsenical encephalopathy, it is concluded that the maintenance of a high level of N.A.B. increases the incidence rate of arsenical encephalopathy. Such an effect of a relatively high arsenic concentration maintained in the body for a comparatively long time indicates a causative relationship between the incidence rate of arsenical encephalopathy and the frequency of injections of N.A.B.

Mechanism

It is generally agreed that the mechanism involved in haemorrhagic encephalopathy is vascular damage;" capillary encephalorrhagy " is the term proposed by Globus and Ginsberg (1933). Also thrombosis has been regarded as the cause of haemorrhagic and necrotic lesions; damage to the endothelium of blood vessels with swelling and fatty degeneration is recorded. The mechanism of perivenous and peri-arterial demyelination is not understood. A toxin has been held responsible by Dorothy Russell. In rare non-haemorrhagic cases oedema and vaso-paralysis (Scheiniker, 1944) have been regarded since Ehrlich (1914) as the cause of the clinical picture and fatal outcome.

An attempt to analyse the lesions observed in this series as part of the mechanism producing arsenical encephalopathy shows the following: the presence of vascular and circulatory disturbances is indicated by stasis, congestion, haemorrhage, and occasional thrombosis; capillary haemorrhages, usually indicating a severe vascular and circulatory disturbance, are scarce.

An important feature of the condition is the presence of perivascular viscus exudation of plasma indicating loss of selective permeability of blood vessels. This is also indicated by the high protein values of the cerebrospinal fluid observed in arsenical encephalopathy, and their importance is stressed by the observation that the average protein content of the cerebrospinal fluid has been significantly lower in cases which recovered than in cases with fatal outcome. It is stressed that this loss of selective permeability recorded here is not associated with gross oedema, the discrepancy between the amount of perivascular viscus exudation and the absence of the effects of severe oedema is obvious. Not only is the intracranial pressure, determined by lumbar puncture, within normal limits, but also the fundus is devoid of papilloedema, and the anatomical manifestations of increased intracranial pressure, that is, pressure cones and flattened gyri, are also absent. It seems of interest that the intracranial pressure was often below average, the blood pressure being normal.

The following working hypothesis may be attempted, although no special claims can be laid on it. Administration of N.A.B. produces damage to cerebral blood vessels, resulting in loss of selective permeability. Subsequently N.A.B. and split products of N.A.B. penetrate into the nervous tissue, mainly perivascularly and peripherally, and interfere there with oxydation systems of the parenchyma. In haemorrhagic cases the vascular damage is more prominent in the form of haemorrhage and oedema; in non-haemorrhagic cases the invisible tissue toxic effect is more prominent and may manifest itself at later stages of the disease as perivascular necrosis and demyelination.

Summary

1. Arsenical encephalopathy in Indian troops is the pathological equivalent of classical haemorrhagic arsenical encephalopathy; it has the same time of onset and the same clinical picture as the classical condition, but it differs by a comparatively high incidence rate and by the absence of gross haemorrhagic or purpuric lesions. The relationship to the classical condition is stressed by occasional typical haemorrhagic cases and by cases showing demyelination.
2. The "racial" incidence of arsenical encephalopathy is high in Indians; it is considered that part of it is "social" incidence.

3. An attack of malaria during or shortly before antisyphilitic treatment is an important factor in increasing the incidence of arsenical encephalopathy.

4. The incidence rate of arsenical encephalopathy in Indian troops treated with weekly injections of N.A.B. is of the same order as the incidence rate in Indian civilians.

5. The increase in the incidence rate of arsenical encephalopathy in Indian troops, as compared with Indian civilians and observed since 1943, is the effect of the treatment of syphilis with bi-weekly injections of N.A.B.

6. The increased incidence rate of arsenical encephalopathy following bi-weekly injections of N.A.B. is not proportionate in all provincial groups of Indian troops. The incidence rate is significantly higher in Madrassis and Maharrattas, and only moderately increased in Indians from other provinces.

7. The incidence rate of arsenical encephalopathy in sero-positive primary and secondary syphilis is not higher than in sero-negative primary syphilis. The higher incidence rate of arsenical encephalopathy in sero-negative primary syphilis as compared with the two other groups is still below statistical significance in this series.

8. Four successive stages of the clinical picture can be separated. The first three stages are reversible, all cases reaching the fourth stage—coma—are fatal.

9. The protein content of the cerebrospinal fluid in arsenical encephalopathy is usually greatly increased without proportionate increase of the cell count; the average protein content in cases which recover is significantly below the average in fatal cases. The high protein content of the cerebrospinal fluid is interpreted as the manifestation of loss of selective permeability of blood vessels.

10. The arsenic content of the brain in this series was relatively high, and of the same order as reported by Osterberg and Kernohan (1934) in cases of classical hemorrhagic arsenical encephalopathy.

11. The tissue pathological findings are signs rather than the cause of the condition. They indicate vascular damage, circulatory disturbance, and loss of selective permeability of blood vessels.

12. Gross edema, increasing the intracranial pressure, is not a feature of the condition. This is substantiated by the absence of pallor, the average pressure of the cerebrospinal fluid at lumbar puncture, and the absence of pressure cones.

13. The fact that a relatively high level of N.A.B. is maintained for a comparatively long time if treatment is given at bi-weekly intervals, together with the subsequently increased incidence rate of arsenical encephalopathy, indicates a causative relationship between the level of arsenic maintained in the tissues, the frequency of injections of N.A.B., and the incidence rate of arsenical encephalopathy.

14. Arsenical encephalopathy subsequent to a standardized form of antisyphilitic treatment is considered here as a tissue toxic effect of trivalent organic arsenical compounds in persons predisposed racially, socially (nutrition), and individually (e.g., malaria). Whereas vascular damage is more prominent in cases of classical hemorrhagic encephalopathy, functional and eventual anatomical damage to the parenchyma was more conspicuous in arsenical encephalopathy in Indian troops.

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