Electrophysiological dysfunction and cellular disruption of sensory neurones during incubation with Treponema pallidum

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SUMMARY Treponema pallidum (Nichols strain) was incubated with cultured nerve cells derived from dorsal root ganglia of rat embryos. The electrophysiological response of these neuronal cells was then investigated. Cells exposed to $2 \times 10^8$ treponemes/ml responded abnormally after 13 hours and failed to respond after 18 hours. In contrast, control preparations exposed to heat-inactivated treponemes or to culture medium responded normally after 72 hours. Extended incubation with viable treponemes resulted in various degrees of nerve cell disruption as shown by scanning electron microscopy. With some cells holes in the cytoplasmic membrane were detected; with others a coagulated matrix of apparent nuclear material and remnants of cytoskeletal elements indicated more severe destruction. These findings may explain the painless nature of many of the clinical manifestations of syphilis as well as the severe damage to central nervous system tissue in tertiary and congenital syphilis.

Introduction

Ovcinnikov and Delektorskij demonstrated treponemal association with nerve cell tissues within material from syphilitic lesions. Although they did not observe nerve histopathology, they suggested that Treponema pallidum might inactivate nerve function; this in turn could explain the painless nature of most of the clinical manifestations of syphilis. The accompanying paper described the interaction of T pallidum with cultured cells derived from dorsal root ganglia, superior cervical ganglia, and spinal cord. After 6 hours' incubation numerous treponemes were attached and the nerve cell morphology was not altered. The purpose of this study was to show that with extended incubation treponemes initially cause electrophysiological dysfunction then morphological disruption of dorsal root ganglia neurones in culture.

Materials and methods

T PALLIDUM

The Nichols strain was maintained by intratesticular passage. Conditions for extraction have been described. After slow-speed centrifugation organisms were centrifuged at high speed $(18,000 \times g$ for 30 minutes at $4^\circ C$) to remove the inflamed testicular constituents. The only departure from previous conditions was the omission of glutathione, cysteine, and dithiothreitol from the treponemal suspending medium. For inactivation by heat treponemes were placed at $56^\circ C$ for 10 minutes.

NERVE TISSUE CULTURE CELLS

Nerve cells were isolated from dorsal root ganglia of 17-19-day-old rat embryos. Cell preparation, medium, and conditions of incubation have been described. After the cells had grown for 3-4 weeks culture medium was removed and the cells were washed three times with 2 ml treponemal suspending medium. Different concentrations of treponemes were then added in a 2-ml volume. Cells were incubated at $37^\circ C$ in an atmosphere of 92-5% nitrogen, 5% carbon dioxide, and 2-5% oxygen.

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SCANNING ELECTRON MICROSCOPY
The techniques and fixation procedures for examining nerve cells using scanning electron microscopy have been described.2

ELECTROPHYSIOLOGICAL METHODS
Electrical recordings were made directly in the culture dishes on the stage of a Zeiss inverted microscope. The tip of a microelectrode was inserted into a nerve cell. Voltage differences were then measured between this intracellular electrode and a second electrode used as a ground immersed in the extracellular fluid. Electrical pulses were passed through the intracellular electrode and the change in voltage across the cell membranes monitored. Microelectrodes were positioned with a Leitz micromanipulator under direct observation at a magnification of × 400. The temperature, monitored by a thermistor, was maintained at 37°C with an overhead heat lamp. Recordings were made in tissue culture medium to avoid disturbances in the infective process and to promote cell survival over an extended period of time.

Fine-tipped glass microelectrodes were filled with 4 mol/l potassium acetate (pH 7.0) with resistances of 30-80 MΩ. A bridge circuit (Mentor N-950, Minneapolis, Minnesota) was used for simultaneous current stimulation and voltage recordings. The cells were stimulated with a depolarising pulse, and the action potentials were monitored on an oscilloscope and recorded on a Hewlett-Packard FM tape recorder. The electrophysiological responses were analysed with a computer programme written for a PDP-12 computer (unpublished data). Values for the different experimental conditions were compared using a standard statistical package.

Results
Neuronal cells were presumptively identified in the previous study2 by their characteristic morphology. Definitive identification is based on the ability of these cells to respond to electrical stimulation with an action potential. Fig 1a shows a typical action potential and defines the indices of resting membrane potential, spike amplitude, maximum rate of rise, and after-hyperpolarisation maximum. Normal action potentials are readily distinguished from abnormal action potentials.

Attempts were made to evaluate the influences of T pallidum on the genesis and character of the action potential of cultured nerve cells. In preliminary experiments control preparations failed to respond appropriately. Analysis of different components of the culture medium showed that glutathione, cysteine, and dithiothreitol prevented typical responses. These three reducing agents were then omitted from the treponemal suspending medium. Various concentrations of treponemes were added to the cultured cells. Controls included culture medium alone or heat-inactivated treponemes. At different times after inoculation neuronal cells were electrically stimulated and the action potentials measured. In the first two experiments the two control preparations showed normal responses after incubation for 72 hours. In contrast, neuronal cells exposed to 2 × 10⁸ treponemes/ml lost their ability to respond after eight hours in one experiment and after 10 hours in the second.

The results of a third experiment are shown in table I. Normal responses were defined as those that fell within ±1 SD of the resting membrane potential and spike amplitude for the controls not containing

**FIG 1(a)** A typical action potential of a dorsal root ganglion neurone. Electrophysiological indices are defined: A, resting membrane potential (mV); B, spike amplitude (mV); C, maximum rate of rise (mV/ms); D, after-hyperpolarisation maximum (mV). (b) Electrophysiological responses of neuronal cells exposed to culture medium or treponemes (A, normal control; B and C, abnormal treponemal-impaired). The calibration scale is 20 mV along the y axis and 1 ms along the x axis.
Treponemal interference with electrophysiological responses of cultured dorsal root ganglia

<table>
<thead>
<tr>
<th>Time (hours)</th>
<th>Normal response* total tested</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without treponemes</td>
<td>2 × 10⁸ Treponemes/ml</td>
</tr>
<tr>
<td>0</td>
<td>5/5</td>
</tr>
<tr>
<td>5</td>
<td>5/5</td>
</tr>
<tr>
<td>7</td>
<td>5/5</td>
</tr>
<tr>
<td>10</td>
<td>5/5</td>
</tr>
<tr>
<td>13</td>
<td>3/11</td>
</tr>
<tr>
<td>18</td>
<td>0/5</td>
</tr>
<tr>
<td>26</td>
<td>10/10</td>
</tr>
</tbody>
</table>

*Defined as those that fall within ± 1 SD of the resting membrane potential and spike amplitude for the controls not containing treponemes.

At 0 hours the nerve cells exposed to control medium and to the two different concentrations of *T. pallidum* functioned normally. After incubation for 26 hours, 10 of 10 cells in the control preparation responded normally. With treponemes at 2 × 10⁸ organisms/ml abnormal responses were detected after 13 hours; only three of 11 cells responded normally. After 18 hours action potentials could not be elicited. With treponemes at 1 × 10⁸ organisms/ml normal responses were detected in five of five cells after 18 hours and in 10 of 12 cells after 26 hours.

Fig 1b shows representative action potentials of control cells (A) and test cells (B and C) incubated with treponemes for 13 hours. The mean values determined by combining the indices for individual nerve action potentials in each experimental condition are shown in table II. For control preparations not exposed to *T. pallidum* no significant differences were apparent after 0 and 26 hours; these two sets of values were combined. Incubation for 13 hours with 2 × 10⁸ organisms/ml resulted in abnormal electrophysiology in eight of 11 neuronal cells. Comparison of the individual indices for these cells showed significant differences in resting membrane potential, spike amplitude, maximum rate of rise, and after-hyperpolarisation maximum compared with the controls. At this time the morphology of the neuronal cells was unchanged.

In further experiments nerve cultured cells were inoculated with 1 to 2 × 10⁸ organisms/ml and controls of culture medium, heated treponemes, and the high-speed supernatant. Cultures were examined with phase-contrast microscopy. After incubation for 24 hours pronounced damage to both neuronal cells and fibroblastic cells was apparent in cultures exposed to viable treponemes. Neuronal morphology was not altered in the control preparations incubated for 24 hours.

Degradation was related to the concentration of *T. pallidum*. Below 7 × 10⁷ organisms/ml no morphological or electrophysiological changes were observed after 48 hours' incubation. After 48 hours, motility of the organisms was sharply decreased probably because of the omission of reducing agents from the culture medium.

Treponemal degradation of neuronal cells was evident by scanning electron microscopy. Cultures were inoculated with 2 × 10⁸ organisms/ml. Figure 2 shows atypical morphology of a nerve cell after 16 hours' incubation. In contrast to the round elevated appearance of normal neurones the cells assumed a flattened morphology, and their surfaces were marked with irregular holes. Some cells were more damaged than others.

Various stages of treponemal-mediated nerve-cell degradation are shown in figs 3-5. Fig 3 shows the ghost-like remnant of an apparent dorsal root ganglia cell. The morphology was greatly altered. The nuclear profile in the centre of the cell deteriorated to a coagulated matrix. Remnants of cellular substratum attachment processes can be seen peripherally. At higher magnification of this peripheral area (fig 4) the edge of the cell is demarcated by fibrous extensions, which are associated with small bleb-like particles. These may

**TABLE II** Selected indices of electrical properties of cultured dorsal root ganglia infected with treponemes

<table>
<thead>
<tr>
<th>Preparation*</th>
<th>Incubation time (hours)</th>
<th>Resting membrane potential (mV)</th>
<th>Spike amplitude (mV)</th>
<th>Maximum rate of rise (mV/ms)</th>
<th>After-hyperpolarisation maximum (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without treponemes (15)</td>
<td>26</td>
<td>55.61 ± 3.85</td>
<td>81.65 ± 9.13</td>
<td>204.11 ± 40.41</td>
<td>6.40 ± 1.48</td>
</tr>
<tr>
<td>With 2 × 10⁸ treponemes/ml (4)</td>
<td>7</td>
<td>54.17 ± 2.40</td>
<td>89.18 ± 3.39</td>
<td>220.60 ± 9.25</td>
<td>3.65 ± 1.02</td>
</tr>
<tr>
<td>(5)</td>
<td>10</td>
<td>54.31 ± 1.25</td>
<td>83.79 ± 5.56</td>
<td>212.34 ± 26.2</td>
<td>4.42 ± 1.18</td>
</tr>
<tr>
<td>(8)</td>
<td>13</td>
<td>50.12 ± 3.86</td>
<td>65.66 ± 16.11</td>
<td>146.64 ± 61.96</td>
<td>2.81 ± 0.99</td>
</tr>
<tr>
<td>With 1 × 10⁸ treponemes/ml (5)</td>
<td>13</td>
<td>56.43 ± 3.45</td>
<td>77.11 ± 8.84</td>
<td>195.40 ± 38.73</td>
<td>4.06 ± 2.11</td>
</tr>
<tr>
<td>(6)</td>
<td>18</td>
<td>54.77 ± 4.50</td>
<td>80.51 ± 14.96</td>
<td>197.34 ± 63.51</td>
<td>6.05 ± 2.02</td>
</tr>
<tr>
<td>(9)</td>
<td>26</td>
<td>50.63 ± 4.29</td>
<td>73.36 ± 14.16</td>
<td>170.52 ± 60.55</td>
<td>5.8 ± 1.73</td>
</tr>
</tbody>
</table>

SEM = Standard error of mean

*No of neuronal cells tested given in parentheses

†Statistically significant, P<0.05
Electrophysiological dysfunction and cellular disruption of sensory neurones by *T. pallidum*

**Discussion**

A number of reports have shown that cultured nerve cells accurately reflect in-vivo phenomenon. These cultured cells elaborate neuritic processes; they respond to voltage stimulation with typical action potentials similar to those generated in vivo; they form functional contacts with one another, and action potentials are transmitted to neighbouring neuronal cells. In addition, different types of nerve tissues that vary in structure and function show the same variations after in-vitro cultivation. Thus, the observations in this paper should be representative of in-vivo phenomenon.

The accompanying study showed that *T. pallidum* attached to cultured cells derived from dorsal root ganglia, superior cervical ganglia, and spinal cord. This study extends these findings to longer periods of incubation and demonstrates treponemal dysfunction and morphological damage to neuronal cells of dorsal root ganglia. This detrimental effect was dependent on attachment of treponemes. Heat-inactivated organisms failed to attach, and correspondingly nerve cell function and morphology were degraded.

**FIG 2** Scanning electron micrograph of a nerve cell derived from dorsal root ganglia incubated for 16 hours with *T. pallidum*. The flattened cell shows irregular holes (arrowheads) on its surface. (× 3900).

represent the cytoskeletal cores of radially extending retraction fibrils.

Further degradation of a nerve cell is shown in fig 5. The cytoplasm is difficult to differentiate from the condensed nucleus. The cell periphery is profiled by small blebs, and the fibrous extensions of the cellular margin have apparently been degraded.
unchanged. Concentrations below $7 \times 10^7$ organisms/ml did not adversely influence nerve cell function or morphology. Damage at this, or lower concentrations, may have resulted if treponemal survival was extended. Without reducing agents, however, organisms retained motility in our system for about 48 hours. Most importantly, the concentrations that were detrimental ($2 \times 10^8$ treponemes/ml) are consistent with syphilitic infection. The critical number of organisms required for the initial manifestations of erythema and induration is approximately $10^7$ treponemes. Assuming a generation time of 30 hours, within 4-5 days and before ulceration occurs, over $10^8$ treponemes should be present. Neuronal cells lost their ability to respond electrophysiologically (8-10 hours) before morphological changes became apparent (16 hours). This is the first report of in-vitro nerve dysfunction and disruption attributed to the attachment of bacteria. A normal action potential requires an intact neuronal membrane. Thus, the first detectable damage must have reflected alterations in nerve cell membranes. With further incubation morphological damage occurred as manifested by holes in the nerve cell membrane and a flattening of the typical raised morphology. Further disruption of nerve cell integrity was characterised by condensation of the nucleus and an outlining of the cellular boundaries by fibrous cytoskeletal remnants. Similar morphological alterations have been shown in cells treated with Triton-X, a detergent that solubilises the plasma membrane, or EGTA, a Ca$^{++}$ specific chelator.
Electrophysiological dysfunction and cellular disruption of sensory neurones by T pallidum

These processes can be easily broken by gentle manipulation and separated from material that remains attached to the substratum. This substrate-attached material mediates cell adhesion and is primarily composed of fibronectin, glycosaminoglycans, and cytoskeletal proteins. These areas of the cultured cells are apparently somewhat more resistant to degradation by T pallidum.

Oakes et al. recently reported similar detrimental influences of herpes viruses on dorsal root ganglia in culture. After viral inoculation alterations in electrophysiological indices were observed within four hours. Morphological alterations were detected after 16 hours; at this time less than half the neuronal cells responded with action potentials. Nerve cells were totally destroyed after 48 hours' incubation. The authors suggested that herpes virus reduced the sodium ion conductance of neuronal cells.

The findings in this study may partially explain two unknown factors of syphilitic infection: (a) the painless nature of most of the clinical manifestations of each stage of infection and (b) the severe nerve degeneration in tertiary and congenital syphilis. Two preliminary reports have shown that treponemes are associated with nerve fibres within lesion material.

In our tissue culture system after treponemal attachment to neuronal cells the initial effect was a functional impairment of the ability of these cells to generate an action potential. With further incubation morphological disruption of the neuronal cells was observed. Clinical manifestations of primary and secondary infection are usually evident for 2-6 weeks,
FIG 5  Nerve cell incubated for 16 hours with T pallidum showing greatly altered morphology. The nuclear and cytoplasmic constituents are difficult to distinguish. The outline of the cell is marked by small cytoplasmic particles, probably remnants of attachment contacts. (× 1400).

whereas manifestations of tertiary syphilis may last for years. In primary and secondary stages nerve deficits do not occur. The presence of treponemes within tissues for shorter periods of time (weeks) may interfere transiently with nerve function and account for the lack of pain. After healing pain receptors would return to normal without nerve deficits. In tertiary and congenital syphilis nerve damage frequently occurs. The presence of treponemes within tissues for longer periods (years) may result in morphological destruction causing damage to the central nervous system, particularly the cerebral cortex, spinal cord, and peripheral nerves.

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