Demonstration of substances capable of contracting smooth muscle in the haematoma fluid from certain cases of ruptured cerebral aneurysm

MONAMY BUCKELL

From the Neurosurgical Research Laboratories, Atkinson Morley's Hospital (Branch of St. Georges Hospital), Wimbledon, London

In a proportion of patients suffering rupture of an intracranial aneurysm, the angiogram demonstrates marked narrowing of the vessels in the proximity of the lesion. These appearances are considered by many neurosurgeons and neuroradiologists to be due to active constriction of the affected vessels.

Others, however, deny the very existence of this phenomenon as far as cerebral vessels are concerned and explain the appearances on the grounds of atherosclerosis, stretching of the vessels by local haematoma or brain swelling, or to inadequate filling with contrast material. To the surgeon who has experience of direct surgical approaches to intracranial aneurysms, the development of marked vasoconstriction is an observable phenomenon: manipulation, for example, of the middle cerebral artery to dissect out an aneurysm will often result in marked shrinkage of the main vessel and its peripheral branches to half their original calibre. This is, of course, a mechanically induced constriction of the vessels.

If the phenomenon is accepted, the causation is not clear, but among the various possibilities, the most popular is that such a change could be produced or maintained by a chemical agent liberated locally in relation to the haemorrhage.

The point is one of importance as there is no doubt that patients with angiograms showing evidence of this vascular narrowing fare worse with surgical intervention when compared with those patients showing vessels of normal calibre.

To test this theory, we examined nine specimens of haematoma material taken from the immediate environment of a ruptured aneurysm. The material was collected at craniotomy undertaken for direct attack upon the aneurysm and thus the most dangerously ill patients were excluded as being unsuitable for this form of treatment (category A cases). In many other patients the quantity of haematoma fluid was too small or was evacuated by suction before a specimen could be obtained.

MATERIAL AND METHOD

The samples consisted of a mixture of old fluid blood with varying amounts of clot, cerebrospinal fluid, and flecks of necrotic brain tissue. Amounts ranging from 0-2 to 2-9 g. were obtained from the immediate neighbourhood of the aneurysm and its parent vessel. In case 1 subdural fluid was also obtained and in case 2 there was a specimen of cerebrospinal fluid that had been taken six days preoperatively and stored frozen. The material was collected with a siliconed syringe or a scoop and transferred to a silicone-coated container. Some specimens reached the laboratory within half an hour; others, taken at another hospital, were transported without refrigeration but with minimal delay, the slowest taking three and a half hours. To avoid activation of protease-peptide systems by contact with glass, which was thought to have occurred in some earlier cases excluded from this series, all handling up to the time of testing was carried out in silicone-coated glassware.

The fluid specimens were centrifuged, or in the case of the mainly solid ones a saline extract was made, and the supernatant fluid was separated. In four cases this was tested at once on a preparation that had been set up in anticipation of a specimen. The other supernatants were frozen immediately and stored at -15°C for the periods shown in the Table.

For the determination of total smooth-muscle-contracting activity and of true 5-hydroxytryptamine (5-HT) con-

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Arterial Narrowing</th>
<th>Days between Haemorrhage and Operation</th>
<th>Days Specimen Stored</th>
<th>Total Activity Expressed as HT Units/specimen</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Yes</td>
<td>3</td>
<td>1</td>
<td>60</td>
</tr>
<tr>
<td>2</td>
<td>Yes</td>
<td>12</td>
<td>1</td>
<td>40</td>
</tr>
<tr>
<td>3</td>
<td>Yes</td>
<td>3</td>
<td>0</td>
<td>88</td>
</tr>
<tr>
<td>4</td>
<td>Yes</td>
<td>3</td>
<td>1</td>
<td>20</td>
</tr>
<tr>
<td>5</td>
<td>No</td>
<td>2</td>
<td>4</td>
<td>25</td>
</tr>
<tr>
<td>6</td>
<td>No</td>
<td>4</td>
<td>0</td>
<td>33</td>
</tr>
<tr>
<td>7</td>
<td>No</td>
<td>2</td>
<td>0</td>
<td>12.5</td>
</tr>
<tr>
<td>8</td>
<td>No</td>
<td>7</td>
<td>2</td>
<td>17.5</td>
</tr>
<tr>
<td>9</td>
<td>No</td>
<td>6</td>
<td>0</td>
<td>64</td>
</tr>
</tbody>
</table>

The Table shows the following results:

1. No. of specimens: 9
2. Days between haemorrhage and operation: 3-12 days
3. Days specimen stored: 1-4 days
4. Total activity expressed as HT units/specimen: 0-88
5. 5-HT activity: not done

tent the rat stomach strip method of Vane (1957) was used, with addition of UML-491 (Methysergide) as a blocking agent for 5-HT. Only preparations capable of detecting one nanogram (ng) of 5-HT in the test dose were used. Where activity was too high to record, dilutions were made in physiological saline. Saline alone had no effect on the preparation but in some specimens dilutions led to an increase of total activity, suggestive of a polypeptide. The total activity of the specimen in terms of standard 5-HT and of acetylcholine was determined, UML-491 was then added to the superfusion fluid to give a concentration of 10⁻⁸. At this concentration UML-491 did not change the sensitivity of the preparation to acetylcholine while it completely inhibited 5-HT. The assay was then repeated in terms of acetylcholine and the proportion of the total activity due to 5-HT was calculated as the decrease of activity relative to acetylcholine after the addition of the 5-HT inhibitor.

RESULTS

The table gives the total activity of each specimen and the actual amount of 5-HT found. The total activity is expressed as equivalents of 5-HT, though not necessarily all due to this substance, 1 'HT unit' being the amount of activity that gave rise to a contraction equivalent to that produced by 1 nanogram of 5-HT. In addition, the subdural fluid from case 1 had a total activity of 22 'HT-units' of which 4 ng. per ml. was 5-HT, and the cerebrospinal fluid from case 2 contained 8 units of total activity per ml. of which 50% was 5-HT.

The finding of 5-HT in cases 1, 3, and 4 and its absence from the four control cases gives a possible chemical explanation for the local state of the vessels. Case 2 did not come to operation until 12 days after the more recent of his two subarachnoid haemorrhages and seven days after the arteriogram. At this time spasm was presumed on clinical grounds still to be present, but the intense spasm seen at the time of arteriography could have lead to infarction. The patient died 11 days post-operatively and an extensive area of cerebral infarctation was seen at necropsy.

Though a high total activity was found no true 5-HT could be demonstrated in the operation specimen from this case. However, the cerebrospinal fluid taken on the day after the arteriogram had one of the highest total activities we have yet found in this material and contained 4 ng. 5-HT per ml.

Raynor, McMurtry, and Pool (1961) showed that topically applied 5-HT could constrict the exposed cerebral vessels of cats, though they failed to obtain a reaction to cerebrospinal fluid from three cases of subarachnoid haemorrhage. We started by examining lumbar cerebrospinal fluid but then decided that the material in contact with the vessels would be more informative and so have been concentrating on specimens removed at operation. The conclusion cannot be drawn that 5-HT is in fact responsible for the arterial narrowing in our patients as there is other material also present in excess in the affected cases; also the chemical differences might be a sequel of the vascular disturbance rather than a cause of it. This small series of results can only point the way to further investigation, but the preliminary findings are reported at this stage as we believe that this is the first demonstration of a chemical difference in the immediate surroundings of affected and unaffected vessels. As a potentially vasoconstrictor substance with known antagonists is involved, the confirmation of its presence by other workers and perhaps by other methods is of importance. Meanwhile we are extending our series of cases and also pursuing the nature of the non-HT activity.

We are grateful to the Wolfson Foundation for financial support, to Drs. J. R. Vane and J. W. Thompson, of the Department of Pharmacology, Royal College of Surgeons, for advice and help with the pharmacological technique, and to Messrs. Sandoz Products Ltd. for a gift of UML-491.

REFERENCES


Demonstration of substances capable of contracting smooth muscle in the haematoma fluid from certain cases of ruptured cerebral aneurysm

Monamy Buckell

*J Neurol Neurosurg Psychiatry* 1964 27: 198-199
doi: 10.1136/jnnp.27.3.198

Updated information and services can be found at: [http://jnnp.bmj.com/content/27/3/198.citation](http://jnnp.bmj.com/content/27/3/198.citation)

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to: [http://group.bmj.com/group/rights-licensing/permissions](http://group.bmj.com/group/rights-licensing/permissions)

To order reprints go to: [http://journals.bmj.com/cgi/reprintform](http://journals.bmj.com/cgi/reprintform)

To subscribe to BMJ go to: [http://group.bmj.com/subscribe/](http://group.bmj.com/subscribe/)