Tranexamic acid (AMCA) in aneurysmal subarachnoid haemorrhage

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Subarachnoid haemorrhage due to a ruptured cerebral aneurysm is a dramatic event in neurosurgery and one of the most destructive. Its management is complicated by the risks of rebleeding and cerebral vasospasm and their sequelae. Rebleeding and vasospasm rarely occur before three days after the primary bleed.\(^1\)\(^,\)\(^2\) Angiographic vasospasm is seen most often 10 to 17 days after bleeding.\(^3\) Because of the brain's vulnerability during spasm many surgeons prefer to wait until two weeks after the primary bleed to obliterate the aneurysm.\(^4\) Sano and Saito\(^5\) reported fatal postoperative vasospasm in patients operated upon on days 4 and 7 after a haemorrhage. However, the 'spasm period' coincides with the time when rebleeding is most likely. The incidence of rebleeding is highest at the end of the first week and at the beginning of the second after the primary bleed, and the mortality after the first recurrence is reported to be between 43\% and 64\%.\(^1\)\(^,\)\(^6\) Thus most surgeons accept spontaneous mortality due to either progressive deterioration or to rebleeding during the first and second week. The surgical mortality when the patient has recovered from this deterioration can be low. It is therefore necessary to look for ways of lessening the risk of rebleeding during the first critical two weeks.

The aim in treating patients with a ruptured aneurysm with antifibrinolytic agents is to prolong the duration of the formed blood clot within and about the wall of the aneurysm and thus prepare the way for mechanical repair of the rupture. Promising results with aminocaproic acid (EACA)\(^7\)\(^-\)\(^11\) and AMCA\(^12\)\(^-\)\(^19\) have been reported. Others, however, found that these drugs had no effect on rebleedings.\(^20\)\(^-\)\(^24\) In this paper I report the results of two controlled clinical trials of the effect of AMCA on aneurysmal rebleeding, vasospasm, hydrocephalus, and circulatory disturbances.

Patients and methods

The trials were conducted over the years 1972 to 1978. The sealed-envelope technique was used instead of the double-blind method, since it was considered unethical to give placebo injections for a prolonged period. Only patients admitted to the hospital within three days after a subarachnoid haemorrhage due to a ruptured aneurysm and in whom treatment was started within 72 hours were included in the trials. The diagnosis was verified by spinal fluid (CSF) examination and cerebral angiography. In the second series all patients were also examined with CT-scan.

The patients were randomly assigned to conservative treatment (bedrest and sedation) or conservative treatment together with the administration of AMCA. In a first series of 46 patients 23 were given AMCA by slow intravenous injection in a dosage of 1 g 4-hourly during the first week, 1 g 6-hourly during the second to fifth weeks inclusive, and 1 g 8-hourly during the sixth week. In a second series of 59 patients 30 received AMCA in hourly intravenous infusions 6 g daily during the first week, 4 g daily during the second week, and 1.5 g by mouth four times a day during the third to sixth weeks. The patients who received AMCA were treated according to this scheme until rebleeding, operation, discharge, or death. Thus not all patients were treated throughout the entire six weeks.

Inotropic drugs and corticosteroids were given in most cases in both groups. The patient's clinical condition on admission was assessed according to Botterell et al.

Evidence of rebleeding was verified by lumbar puncture, CSF spectrophotometry, CT-scan, or angiography. Repeated preoperative CT-scans and echo-encephalograms were performed to detect possible haematoma and hydrocephalus. The incidence of vasospasm was observed by repeated angiograms. The diameters of the internal carotid, middle cerebral, and anterior cerebral arteries were measured and compared with the diameters of the same vessels on the admission angiograms.

Results

First series

Twenty-three of the 46 patients were controls and
Twenty-nine of the 59 patients in this series were controls and 30 were treated with AMCA. The groups matched for sex and condition on admission. The mean age for men was 44 years (range 27 to 72) and for women 55 years (range 19 to 73). Hyper-tension was noted in seven AMCA-treated and 12 control patients. In the AMCA-treated group a total of 14 patients were operated on at an average of 14 days after the primary bleed, whereas 22 patients in the control group were operated on on average 16 days after bleeding.

In the AMCA-treated group there were six rebleedings in six patients occurring 7 to 24 days after initial bleed. Five of them died. In the control group seven patients rebled 3 to 19 days after the initial bleed. Two patients rebled twice and one rebled three times. Five of the patients died from their recurrent bleedings. There was thus a higher rebleeding rate among the AMCA-treated patients in the second series as compared with the first one. This leads us to consider the way of administering AMCA, since the one difference between the two series was the route by which AMCA was given. In the first series it was given by slow intravenous injection for the entire six weeks, and in the second series it was given by intravenous infusion for the first two weeks and then by mouth.

To find out the comparative uptake of AMCA when given by intravenous injection, infusion, or mouth we studied the concentration of the drug and fibrinolytic split products in the CSF in six patients, two of whom received the drug intravenously by injection, two by infusion, and two by mouth. The CSF concentration of AMCA reached 2 mg/l (which was considered therapeutic) within 48 hours in all six patients (Figure). Concomitantly CSF fibrinolytic split products fell regardless of the route of AMCA administration. We concluded, therefore, that the method of giving

**Fig. Concentrations of fibrinolytic split products (FDP) and tranexamic acid (AMCA) in cerebrospinal fluid (CSF) after giving AMCA intravenously or by mouth. After 48 hours concentration of AMCA is 2 mg/l and that of FDP unmeasurable regardless of route of administration of AMCA. SAH = Subarachnoid haemorrhage.**
the drug probably did not affect the rebleeding rate.

Four additional AMCA-treated patients and two controls died of cerebral ischaemia. The majority of patients in each group were examined repeatedly with preoperative angiograms and 19 of the treated and 17 of the control patients showed spasm. Two of the treated patients showed spontaneous aneurysmal thrombosis during treatment. Both had pronounced angiographic spasm. Three treated and seven control patients had preoperative ventricular dilatation. After three to 41 months eight treated patients and 12 controls had slight to moderate ventricular dilatation. Two of the treated patients required a shunt. Deep venous thrombosis in the legs developed in two treated patients and three controls, whereas four treated and one control patient had pulmonary embolism. Four treated and one control patient developed myocardial infarction. Thus thromboembolic complications occurred more often in the AMCA-treated patients. The mean follow-up period was 25 months (range 2 to 41). By then a total of 22 patients had died—13 treated and 9 control patients. The morbidity was similar in the two groups.

**Discussion**

Six controlled clinical trials of the use of AMCA in subarachnoid haemorrhage have been reported (see Table 1). Three reported that AMCA had a positive effect in preventing rebleeding, whereas the others reported practically no difference in the rebleeding rate between AMCA-treated patients and controls. Nevertheless, some factors might have influenced the latter results. Firstly, van Rossum and coworkers did a multicentre study, and in some of the patients treatment started late. Only 25 out of 51 patients underwent cerebral angiography, which leaves doubt about the diagnosis of ruptured aneurysm. Furthermore, three patients had diabetes mellitus. Glucose increases blood fibrinolytic activity and antidiabetic drugs influence the fibrinolytic activity of blood. One of van Rossum's patients had nephritis, and AMCA should not be given to patients with impaired renal function.

Finally, four of his patients were receiving anticoagulants, which could also have influenced the rebleeding rate. In Kaste and Ramsay's series nine patients did not have their rebleedings confirmed by lumbar puncture or neuroradiology. It is difficult clinically to differentiate between rebleeding and vasospasm. This could therefore be a source of error in Kaste and Ramsay's series. The total rate of rebleeding in all six series was thus 19% with 11% mortality among 156 treated patients and 30% rebleedings with 18% mortality among the 153 control patients.

Despite my comments on van Rossum's and Kaste and Ramsay's series, Table 1 shows a clear reduction in rebleedings among AMCA-treated patients whereas the difference in mortality from rebleeding between AMCA-treated patients and controls is rather small. If we compare the incidence of rebleedings in our two series there is an astonishing difference between the one out of 23 AMCA-treated patients and the nine out of 23 controls in the first series and the corresponding six out of 30 and seven out of 29 in the second series (Table 2).

A total of seven patients (13%) in the AMCA-treated group had seven rebleedings at the 16th day on average after the initial haemorrhage, whereas 16 control patients (31%) had 22 rebleedings on the 10th day on average. The difference in rebleeding rate between the two groups is statistically significant (p < 0.05). Most of the recurrent bleedings in the control group occurred during the first week—earlier than in the AMCA-treated group. A total of six treated patients and eight controls died from their rebleedings. In addition, seven treated patients and two controls died from cerebral ischaemia. The total mortality from rebleeding and spasm/ischaemia was 25% in the AMCA-treated group and 19% in the control group during the six weeks' observation (Table 2). The morbidity was the same in both groups.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Rebleedings and subsequent deaths in controlled clinical trials of AMCA in the treatment of subarachnoid haemorrhage</th>
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<tbody>
<tr>
<td></td>
<td>AMCA</td>
</tr>
<tr>
<td></td>
<td>Rebleedings</td>
</tr>
<tr>
<td>Fodstad et al</td>
<td>1/23</td>
</tr>
<tr>
<td>Van Rossum et al</td>
<td>5/26</td>
</tr>
<tr>
<td>Chandra</td>
<td>1/20</td>
</tr>
<tr>
<td>Fodstad et al</td>
<td>6/30</td>
</tr>
<tr>
<td>Total</td>
<td>30/156 (19%)</td>
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</tbody>
</table>
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Table 2  Total mortality from rebleeding or cerebral ischaemia in controlled clinical trials of AMCA in the treatment of two series of patients\textsuperscript{15} 16 32 with subarachnoid haemorrhage over a period of six weeks

<table>
<thead>
<tr>
<th></th>
<th>AMCA</th>
<th>Controls</th>
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</thead>
<tbody>
<tr>
<td>Series 1\textsuperscript{15} 16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deaths from rebleeding</td>
<td>1/23</td>
<td>3/23</td>
</tr>
<tr>
<td>Deaths from ischaemia</td>
<td>2/23</td>
<td>0/23</td>
</tr>
<tr>
<td>Total</td>
<td>3/23</td>
<td>3/23</td>
</tr>
<tr>
<td>Series 2\textsuperscript{24}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deaths from rebleeding</td>
<td>5/30</td>
<td>5/29</td>
</tr>
<tr>
<td>Deaths from ischaemia</td>
<td>5/30</td>
<td>2/29</td>
</tr>
<tr>
<td>Total</td>
<td>10/30</td>
<td>7/29</td>
</tr>
<tr>
<td>Both series</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>13/53 (25%)</td>
<td>10/52 (19%)</td>
</tr>
</tbody>
</table>

In a co-operative aneurysm study Nibbelink\textsuperscript{8} concluded that antifibrinolytic therapy combined with antihypertensive medication caused a higher rate of rebleeding and mortality than antifibrinolytic therapy alone. In our series, one AMCA-treated patient who had a rebleed was taking chlorpromazine, which prolongs the bleeding time\textsuperscript{40} and increases blood noradrenaline.\textsuperscript{41} This in turn might increase the fibrinolytic activity of blood.\textsuperscript{42} 43 Other drugs that might have influenced fibrinolysis-coagulation and thus the rebleeding rate in our patients were corticosteroids\textsuperscript{44} 45 and radiographic contrast media.\textsuperscript{46} The raised intra-arterial pressure during injection of contrast material could also have initiated the aneurysmal rupture in one AMCA-treated patient who rebled during control angiography on the 6th day of treatment.\textsuperscript{47}

However, we have no satisfactory explanation for the difference in the incidence of rebleeding between the two AMCA-treated series, nor do we know why there was a high incidence of cerebral ischaemia among AMCA-treated patients. From their experience of a single or a limited number of cases several authors have expressed concern about an increased incidence of thrombotic and cerebral ischaemic complications associated with antifibrinolytic drugs.\textsuperscript{48} 55 This impression is confirmed by our findings. The sympathomimetic property of AMCA\textsuperscript{55} causing vasospasm and reducing cerebral blood flow must be considered. Earlier experimental work\textsuperscript{54} 56 showed that a direct toxic effect of antifibrinolytic agents on the arterial vessel wall could be of importance. Finally, there is the possibility of interaction with other drugs and substances causing secondary vessel wall changes and circulatory disturbances.

We cannot comment on the incidence of other circulatory complications. Theoretically antifibrinolytic drugs may prevent the dissolution of clots in the subarachnoid space and promote basal arachnoiditis with secondary hydrocephalus. We found more adhesions around the aneurysms at operation in AMCA-treated patients. Knibestol \textit{et al}\textsuperscript{57} found ventricular dilatation in 63\% of 93 AMCA-treated patients and 39\% of 51 controls. In a later follow-up examination, however, there was no difference between the groups. Park\textsuperscript{41} found a 43\% incidence of ventricular dilatation in 46 patients treated with EACA against 17\% in 48 controls. We did not see any more persistent ventricular dilatations in patients treated with AMCA.

The results of our studies support the hypothesis that the risk of rebleeding in patients with haemorrhage due to ruptured cerebral aneurysm is less when they are treated with AMCA during the first two weeks. Nevertheless, AMCA seems to produce cerebral ischaemic complications. This leaves us with the important question: Is it justified to give antifibrinolytic drugs in the preoperative management of patients with subarachnoid haemorrhage? On the basis of our two series we suggest that AMCA be given for only one to two weeks after the primary bleed. Its value in preventing rebleeding must be weighed against its possible side effects.

This investigation was approved by the Ethical Committee at the Umeå University Hospital and by the Swedish National Board of Health and Welfare, Department of Drugs.

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