Short report

Outcome from “rescue clipping” of ruptured intracranial aneurysms during induction anaesthesia and endotracheal intubation

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SUMMARY Eight patients with ruptured aneurysms during induction anaesthesia and endotracheal intubation underwent an emergency “rescue clipping” of their lesion. Three patients died. Of the five survivors three made a good final recovery, one patient was moderately disabled and one remained in coma. Conservative management of this crisis is doomed to failure but the comparatively good outcome of the operative cases supports emergency “rescue clipping”.

There appear to be no published reports on the nature and significance of haemorrhage from a ruptured aneurysm during induction anaesthesia and endotracheal intubation. Once it has been recognised, the patient is either returned to the ward on supportive conservative treatment and, should he survive, another attempt at elective surgery is then undertaken, or the operation is continued. Past experience indicates that under these circumstances patients almost invariably die. The outcome is reported of eight patients who had “rescue clipping” of aneurysms which ruptured during induction anaesthesia and endotracheal intubation.

Material and methods

(a) Cases

Eight patients of a total of 404 (2%) operation inductions for aneurysmal surgery rebled during induction and immediately thereafter underwent an emergency “rescue clipping” of their aneurysm. There were six females and two males; with ages between 33 to 60 years. All patients had previously had uneventful retrograde right brachial angiography or direct carotid angiography under endotracheal general anaesthesia.

(b) Anaesthetic details

The patients studied were evenly distributed amongst four anaesthetists with varied anaesthetic techniques. Althesin, suxamethonium and thiopentone were used for induction and maintained with fentanyl, droperidol, allagaran, etomidate, N2O/O2 mixture in different combinations. Comments such as “difficult”, “awkward intubation”, “coughed during intubation” were made on three patients. In two other adult patients a small (No 8 Portex) size endotracheal tube was used after failure to pass a standard No 9. In two further patients relaxation was achieved only after a second dose of suxamethonium but in the eighth patient there was no indication of difficult intubation. Oropharyngeal local anaesthesia to suppress vagal stimulation during intubation was not used in any of our patients. Neither alpha nor beta adrenergic blocking agents were given prior to or during intubation and induction anaesthesia.

(c) Per-operative details

Blood pressure was measured directly via a radial catheter using a Hewlett-Packard transducer and monitor throughout the operation. The following observations were generally made in all patients. During the scalp incision the surgeon commented on excessive blood ooze although intra-arterial pressure during this period showed no elevation. After turning the flap, however, the initial suspicion of rebleeding was confirmed. The dura and underlying brain were extremely tight with subdural fresh blood and active arterial bleeding from the aneurysm site. Brain swelling was little relieved by ventricular puncture or infusion of 20% mannitol. Only when the blood pressure was reduced to levels of 60 mm Hg systolic level, was swelling reduced sufficiently to allow the surgeon to retract the
Outcome from "rescue clipping" of ruptured intracranial aneurysms

Table

<table>
<thead>
<tr>
<th>Age-Sex (yr)</th>
<th>Site of aneurysm</th>
<th>Days elapsed (from haemorrhage to surgery)</th>
<th>Management data (*)</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Pre-operative</td>
<td>Post-operative</td>
</tr>
<tr>
<td>45, F</td>
<td>Bifurcation middle cerebral</td>
<td>9</td>
<td>C</td>
<td>B, D, E, G</td>
</tr>
<tr>
<td>38, M</td>
<td>Bifurcation middle cerebral</td>
<td>8</td>
<td>A, D</td>
<td>A, B, E, I, J</td>
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<tr>
<td>60, F</td>
<td>Pericallosal</td>
<td>9</td>
<td>C, D, E, I</td>
<td>B, G, H</td>
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<tr>
<td>58, F</td>
<td>Anterior communicating</td>
<td>7</td>
<td>C</td>
<td>B, E, G, H, I</td>
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<td>Posterior communicating</td>
<td>32</td>
<td>D, G</td>
<td>B, G</td>
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<tr>
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<td>Posterior communicating</td>
<td>6</td>
<td>C, D, E</td>
<td>B, E, G, J</td>
</tr>
<tr>
<td>33, F</td>
<td>Bifurcation middle cerebral</td>
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<td>A, C, D</td>
<td>E, J</td>
</tr>
<tr>
<td>35, F</td>
<td>Anterior communicating</td>
<td>7</td>
<td>D</td>
<td>A, B, E</td>
</tr>
</tbody>
</table>

(*) Management data

A Sedatives, tranquilizers
B Anticonvulsants
C Narcotics, analgesics
D Antifibrinolytics
E Steroids
G Mannitol
H Dextran
I Ventricular CSF drainage
J ICP monitoring

GOS (Glasgow Outcome Scale)

1. Good recovery: Patient can lead a full and independent life with or without minimal neurological deficit.
2. Moderately disabled: Patient having neurological or intellectual impairment but is independent.
3. Severely disabled: Conscious patient but totally dependent on others to get through the activities of the day.
4. Vegetative survival.
5. Dead.

brain and clip the aneurysm neck in the usual manner. In one patient (CT, 35, F; table) hypotension could not be induced and partial frontal lobe resection was necessary to achieve exposure. The outcome at the end of a 2 year follow-up period was assessed on the Glasgow Outcome Scale.  

(d) Immediate post-operative management

All patients were able to breathe spontaneously and deliberate mechanical ventilation was not used. The patients remained in the intensive care unit for a few days until their condition was stable and satisfactory.

Results

All patients were grade I to II (Botterell classification) immediately before operation. Immediately after operation all eight patients were breathing spontaneously. Three patients were stuporous and one was mute, all with severe focal deficits including cranial nerve involvement; of the remaining four patients who regained consciousness only one (BH, 42, F) was left with focal neurological deficits.

The table lists aneurysm site, time elapsed from ictus to surgery, pre- and post-operative supportive conservative treatment and the final outcome and date of latest review. Three patients died (mortality rate 37-5%). Of the survivors (62-5%) three (37-5%) made a good recovery at the end of 12 to 18 months; one (12-5%) was moderately disabled and the other remained comatose.

Discussion

The risk of spontaneous recurrent haemorrhage after subarachnoid haemorrhage varies from 14-30% being highest between the 7th to 14th day from ictus.  A The peak incidence of recurrent haemorrhage coincides with lysis of blood clot by fibrinolytic activity but there is the additional risk of recurrent haemorrhage particularly during angiography as well as induction anaesthesia and endotracheal intubation.

Angiographic procedures may be responsible for aneurysmal rupture and subsequent leakage of contrast material into the surrounding subarachnoid spaces and tissues.  A The prognosis from ruptured aneurysm during angiography is very poor and of the 31 cases reported to date 27 (75%) died. Only two (6-4%) of the remaining seven survivors made a complete recovery whereas five developed disability.

Tachycardia and hypertension are well documented complications of laryngoscopy and tracheal intubation in normotensive patients under a variety of anaesthetic techniques similar to the ones used in this study. These hypertensive responses of normal subjects might be enhanced and prove dangerous to hypertensive subjects during intubation, even if the latter are under well controlled antihypertensive therapy.  B The precise cause of the cardiovascular changes following intubation is uncertain. It could, however, be adequately explained on the basis of a reflex sympathetic response to the mechanical stimulation of larynx and trachea. Contributory factors of hypertension and tachycardia may well be due to a continued preoperative anxiety or reflex baroreceptor effect due to minor hypotension after thiopentone and possibly due to suxamethonium.  C These factors appear to
be less important than local laryngotraheal stimulation, particularly since the maximum rise in blood pressure follows endotracheal stimulation. Moreover, it has been shown in cats that the hypertensive response during intubation is due to increased activity of the cardiac sympathetic nerves and that the pattern of the cardiovascular changes was almost identical to man's.15

Although transient arterial hypertension and tachycardia are probably unimportant in most patients, in certain patients myocardial ischaemia,16 left ventricular failure14 and cerebral haemorrhage,17 may occur. Attempts to control these responses have included the administration of various adrenergic blocking agents systemically as well as oropharyngeal local anaesthesia.18 Both alpha19,20 and beta blocking agents20,21 have been successfully used in suppressing the cardiovascular changes during intubation and induction anaesthesia. The effectiveness of adrenergic blocking therapy during induction and intubation of cases with ruptured aneurysm seems worth further study.

The prognosis of recurrent haemorrhage during pre-operative induction is better than that for angiography. The incidence of operative induction haemorrhage is difficult to estimate because the condition has not been reported as such. Girvin22 noted incidentally that three of 66 patients (4.6%) receiving tranexamic acid had operative induction haemorrhage. Immediate operation and evacuation of clot or decompression may be of crucial importance by relieving the enormous rise of intracranial pressure. Massive increase in intracranial pressure at the time of haemorrhage is well documented.23 There was indeed every indication of raised intracranial pressure during the emergency operation with excessive scalp bleeding, dural and brain "tightness", and no response to anti-oedema agents. Hypotension, however, and resultant hypoperfusion produces improvement suggesting that brain swelling rather than oedema is the cause. Mannitol and steroids even in large quantities therefore are ineffective in reducing intracranial hypertension. The immediate reduction of intracranial hypertension prevents or eliminates further mechanical and hypoxic brain damage and prevents the transient physiological lesion from becoming irreversible. Such an injury is manifest by the patient's neurological condition and in particular his level of consciousness. The post-operative state of consciousness is of great prognostic importance. Patients who recovered consciousness immediately after the procedure had a very good to excellent recovery. In contrast those who failed to regain consciousness either died or remained in a vegetative state. Conservative management of this crisis is doomed to failure but the excellent outcome in the operated cases recommended emergency "rescue clipping" of ruptured aneurysms during induction anaesthesia and endotracheal intubation. We would emphasise, however, that this is only possible by prompt and skilful anaesthetic management.

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References
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