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Complications and outcome in patients with aneurysmal subarachnoid haemorrhage: a prospective hospital based cohort study in The Netherlands

Y B W E M Roos, R J de Haan, L F M Beenen, R J M Groen, K W Albrecht, M Vermeulen

Abstract

Objective—The aim of this study was to investigate prospectively in an unselected series of patients with an aneurysmal subarachnoid haemorrhage what at present the complications are, what the outcome is, how many of these patients have “modern treatment”—that is, early obliteration of the aneurysm and treatment with calcium antagonists—what factors cause a delay in surgical or endovascular treatment, and what the estimated effect on outcome will be of improved treatment.

Methods—A prospective, observational cohort study of all patients with aneurysmal subarachnoid haemorrhage in the hospitals of a specified region in The Netherlands. The condition on admission, diagnostic procedures, and treatments were recorded. If a patient had a clinical deterioration, the change in Glasgow coma score (GCS), the presence of focal neurological signs, the results of additional investigations, and the final diagnosis cause of the deterioration were recorded.

Clinical outcome was assessed with the Glasgow outcome scale (GOS) at 3 month follow up. In patients with poor outcome at follow up, the cause was diagnosed.

Results—Of the 110 patients, 47 (43%) had a poor outcome. Cerebral ischaemia, 31 patients (28%), was the most often occurring complication. Major causes of poor outcome were the effects of the initial haemorrhage and rebleeding in 34% and 30% of the patients with poor outcome respectively. Of all patients 102 (93%) were treated with calcium antagonists and 45 (41%) patients had early treatment to obliterate the aneurysm. The major causes of delay of treatment were a poor condition on admission or deterioration shortly after admission, in 31% and 23% respectively.

Conclusions—In two thirds of the patients with poor outcome the causes of poor outcome are the effects of the initial bleeding and rebleeding. Improved treatment of delayed or postoperative ischaemia will have only minor effects on the outcome of patients with subarachnoid haemorrhage.

Keywords: aneurysmal subarachnoid haemorrhage; timing of surgery; complications

Reviews on aneurysmal subarachnoid haemorrhage suggest that with modern treatment strategies, cerebral ischaemia is the only complication that has remained as a cause of poor outcome. The modern treatment includes treatment with calcium antagonists, such as nimodipine, to prevent cerebral ischaemia, and by far the most important, the prevention of rebleeding by occlusion of the aneurysm within 3 days after the initial bleeding.

In a previous study, however, we found that in neurosurgical centres using modern treatment strategies, despite the aim to prevent rebleeding by early surgery, 45% of the patients was not operated on within 3 days after the initial aneurysmal subarachnoid haemorrhage and that rebleeding was still the major cause of poor outcome. Similar results were found in two recently published population based studies in King County, USA and in greater Cincinnati, USA. None of these studies, however, investigated which factors caused the delay in surgery.

The aim of the present study was therefore to reconfirm in an unselected regional cohort of patients with aneurysmal subarachnoid haemorrhage our previous results concerning the timing of surgery and to investigate prospectively which factors are actually responsible for the delay in surgery. Additionally, we investigated how many patients are treated nowadays with calcium antagonists to prevent cerebral ischaemia, what the complications after aneurysmal subarachnoid haemorrhage are at present, and what the outcome of patients is in such a series. Finally, we estimated what the effect might be of improved treatment of complications after subarachnoid haemorrhage on the final outcome.

Patients and methods

STUDY DESIGN

The study design was a prospective cohort study of consecutively admitted patients with aneurysmal subarachnoid haemorrhage in all hospitals in two regions of the Netherlands, Noord-Holland and Flevoland (appendix). According to official figures from the Dutch Central Bureau of Statistics (CBS), a governmental institution, this region has about 2 000 000
inhabitants. All 12 regional hospitals in this region refer their patients with aneurysmal subarachnoid haemorrhage for neurosurgical inter-
ventions to the three neurosurgical units in Amsterdam. These neurosurgical units are all situated less than 85 km (50 miles) from the most remote regional hospital. All three neuro-
surgical units adhere to the same management protocol which includes treatment with the cal-
cium antagonist nimodipine (2 mg/hour intrave-
nously or 6×60 mg/day orally), hypervolaemia, hypertensive treatment to prevent delayed cer-
ebral ischaemia (minimum of 3 l fluid intake daily), and early surgery to prevent rebleeding. The three neurosurgical units in Amsterdam are all teaching hospitals and within
each centre at least three neurosurgeons operate on cerebral aneurysms.

PATIENT SELECTION
All patients with the diagnosis of aneurysmal subarachnoid haemorrhage, primarily admit-
ted in the participating centres, were prospectively studied during a 1 year period (February
1996-February 1997). The diagnosis was based on clinical signs and symptoms and an
aneurysmal bleeding pattern on the initial CT. If the initial CT was negative and subse-
quent CSF examination positive (xanthochro-
ia confirmed by spectrophotometry) an
aneurysm had to be confirmed by angiography
before the patient could be included. Patients
with a perimesencephalic bleeding pattern or
another non-aneurysmal pattern of haemor-
rhage on CT were excluded.

DATA COLLECTION AND MANAGEMENT
Data registration was done in the participating
centres by two research nurses on two portable
personal computers equipped with an elec-
tronic case record form/database program spe-
cially developed for this study. This program
generated electronic forms in which all pa-
tients’ data were directly recorded.

Registration of baseline and clinical data
started on admission in the hospital where the
patient was seen initially. The following base-
line characteristics were recorded: age; sex;
date of aneurysmal subarachnoid haemor-
rhage, and date of admission. Clinical details
recorded at baseline were the Glasgow coma
scale (GCS) and the grade of subarachnoid haemorrhage (World Federation of Neurologi-
cal Surgeons (WFNS)) score on admission, the
results of CT, CSF examination, and the
location(s) of (an) aneurysm(s) on angiog-
raphy.

After admission patients were followed up on
da day to day basis during their whole period in
hospital. All details on treatments used and
management strategies (such as fluid manage-
ment, drug therapies, operative interventions,
consulted specialists, and transfers to other
hospitals or other wards such as intensive care)
and all additional diagnostic procedures (like
blood samples, radiological investigations)
were recorded.

Every time a patient had a clinical deteriora-
tion an additional form was filled out: the date
of the deterioration, the change in GCS, the
onset of the deterioration (sudden versus
gradually), the presence of focal neurological
signs, and the final diagnosed cause of the
deterioration after appropriate investigations
were recorded. The final assessment of pa-
tient’s complication was performed by two of
us (YR and MV), using clinical data, CT data,
and necropsy results. Rebleeding and delayed
cerebral ischaemia were defined as previously
described as definite rebleeding or probable
rebleeding and definite infarction or probable
infarction. Hydrocephalus was defined as the
gradual deterioration of consciousness with
CT evidence of hydrocephalus and no other
explanation for deterioration. Postoperative
ischaemia was defined as a change in the level
of consciousness or the development of focal
neurological signs noticed immediately after
recovery from anaesthesia compared with the
preoperative status, with no evidence of
rebleeding or hydrocephalus on CT or at
necropsy.

Functional health outcome was assessed with
the Glasgow outcome scale (GOS). The
GOS is an index score ranging from 1 (death)
to 5 (good recovery). Death, persistent vegeta-
tive state, and severe disability on the GOS
were combined as poor outcome and moderate
disability and good recovery were recorded on
the study forms as good outcome. In patients
with poor outcome at 3 month follow up, one
of us (YR) reviewed the medical record of the
patient to determine and categorise the actual
cause for this poor outcome in five major
categories: initial bleed; rebleeding; cerebral
ischaemia; operative complications or other
(non-neurological) complications, which had
to be specified. Poor outcome caused by the
effect of the initial bleeding was defined as
impaired consciousness or focal neurological
signs existing since the initial bleed, without
signs of rebleeding, ischaemia, or hydrocepha-
lus on CT or at necropsy.

ETHICAL CONSIDERATIONS
The study protocol was approved by the local
legal advisor or ethics committee of each
participating centre. Informed consent was
asked for in all patients (or legal representative)
with the exception of those who died shortly
after admission, in which case there was no
need for active participation of the patients or
their relatives.

During the whole study period patients were
identified by a unique patient identification
number (PIN) to protect privacy. The code key
to this PIN number remained the responsibil-
ty of the treating physician of the hospital where
the patient was first seen during the study
period and was destroyed after completion of
the whole data set.

Results
During the 1 year study period 110 patients
with an aneurysmal subarachnoid haemor-
rhage were admitted to the participating
regional hospitals and neurosurgical centres.
Table 1 shows the baseline characteristics of
these patients, which are comparable with
baseline characteristics in other population


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In the group of 18 patients with a rebleed, seven had this complication within 2 days after the initial haemorrhage, four on the day of the haemorrhage. Five rebleeds occurred in patients in whom treatment of the aneurysm had been postponed. In three patients with a rebleed, the initial bleed had not been recognised as an aneurysmal subarachnoid haemorrhage. Two patients had a rebleed during surgery and one after surgery. One patient had a rebleed after a negative angiography. Ten of the 18 patients who had a rebleed were in a good clinical condition (WFNS 1 or 2) when surgery was considered. In this group of "good grade" patients, four patients had the rebleed the day after their SAH. Another seven patients had coiling of their aneurysm after a delay of at least 3 days. In these patients the aneurysm was firstly considered unsuitable for surgery; thereafter the question was whether coiling was an option. This led to delayed coiling due to delayed consultation. Surgical clipping was the preferred treatment to occlude the aneurysm in 66 patients (60%). However, in 24 of these patients (36% of the patients with surgery) surgery was postponed for more than 3 days after aneurysmal subarachnoid haemorrhage and 34 of all patients (31%) had no surgery or coiling of the aneurysm.

In 102 (93%) of the patients treatment with calcium antagonists was started. In seven of the remaining eight patients calcium antagonists were not given because these patients were in a poor clinical condition on admission; all died within 3 days after the initial bleeding.

Table 5 lists the causes of delay of treatment to occlude the aneurysm. Deterioration after admission included rebleeding, ischaemia, and hydrocephalus. Delay in referral to the neurosurgical centre was caused by a delay in recognition of the aneurysmal subarachnoid haemorrhage before admission in five patients and in four patients after admission to the regional hospital. Of the eight patients with a negative angiogram, in two an aneurysm was demonstrated after angiography had been repeated. In 10 patients the aneurysm was considered difficult to operate on and in seven patients coiling was preferred. The planning of this procedure led to delayed treatment.

Inclusion period (February 1996-February 1997) coiling of aneurysms in the acute phase just started in this region of The Netherlands. Three patients had coils placed within 3 days after their SAH. Another seven patients had coiling of their aneurysm after a delay of at least 3 days. In these patients the aneurysm was firstly considered unsuitable for surgery; thereafter the question was whether coiling was an option. This led to delayed coiling due to delayed consultation. Surgical clipping was the preferred treatment to occlude the aneurysm in 66 patients (60%). However, in 24 of these patients (36% of the patients with surgery) surgery was postponed for more than 3 days after aneurysmal subarachnoid haemorrhage and 34 of all patients (31%) had no surgery or coiling of the aneurysm.

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Discussion
The incidence of aneurysmal subarachnoid hemorrhage found in this study, 5.5/100 000 a year, is not much different from estimates of this incidence in the Dutch population. Apparently, few patients die before reaching hospital and in a few patients aneurysmal subarachnoid hemorrhage is not recognised. Similarly, in a population based study from King County, Washington, USA only 3% of patients with aneurysmal subarachnoid hemorrhage had died before admission. Therefore, the results of this study are probably representative for all patients with aneurysmal subarachnoid hemorrhage.

The most often occurring complication is cerebral ischaemia, but the proportion of patients with a rebleed is still 16%. However, cerebral ischaemia is not the major cause of poor outcome as the effects of the initial bleed and rebleeding account for two thirds of all poor outcomes.

This study shows that despite aiming at modern treatment strategies the proportion of patients with poor outcome is still 43%. This percentage is much higher than in other studies but these studies have usually been carried out in neurosurgical centres and often consist of randomised clinical trials. Therefore, these figures come from highly selected groups of patients.

Modern treatment consists of early obliteration of the aneurysm and administration of calcium antagonists. Calcium antagonists were given to most of the patients (93%) but early treatment of the aneurysm was carried out in less than half.

We expected that logistic reasons such as, for instance, a delay in angiography at the department of radiology or difficulties in finding an available theatre or a surgeon on short notice would be the major causes of delay. Logistic reasons however, seemed to play hardly any part. The major causes of delaying treatment were the condition of the patient on admission and deterioration from complications such as rebleeding and ischaemia occurring before the planned surgery or coiling to occlude the aneurysm.

The proportion of patients with a rebleed is still high but not all these rebleeds can be prevented by early treatment of the aneurysm. In the group of 18 patients with a rebleeding, in seven this occurred within 2 days of the initial haemorrhage and in two during surgery. Very rigorous treatment to occlude the aneurysm as early as possible in all patients, irrespective of the clinical condition on admission, may half the number of patients with rebleeding. As rebleeding is the cause of poor outcome in about 30% of the patients, this management would lead to a reduction of poor outcome in about 15%.

Similarly, if it is possible to reduce the occurrence of cerebral ischaemia by 50% by a new treatment, irrespective of whether ischaemia developed unrelated to surgery or during or immediately after surgery, the reduction of poor outcome will probably be less than 15%. However, these reductions are difficult to achieve. This study shows that treatment to prevent rebleeding is usually postponed because of the poor clinical condition of the patient on admission or because of deterioration shortly after admission. The results of surgery in this group of patients are less good than in patients who undergo surgery in a better condition. Moreover, in this group of patients outcome may be poor despite prevention of rebleeding, therefore the impact of surgery on outcome will probably be much less than the above estimated 15%. In this study only three of the 18 rebleeds could have been prevented if surgery had been performed in all good grade patients on the same day that the aneurysm was demonstrated. It might be that better results can be achieved by using endovascular techniques to occlude the aneurysm in the acute phase, but also with this treatment the devastating effects of the initial bleed or early complications will remain.

A new effective treatment for cerebral ischaemia might have a different impact in the different types of cerebral ischaemia, as the mechanism of postoperative ischaemic damage is probably not the same as in delayed cerebral ischaemia. If treatment works only on delayed cerebral ischaemia and is very effective by reducing half of the occurrences of this type of ischaemia, the reduction of poor outcome will not be more than 10%. If the effectiveness is less—for instance a 30% reduction of cerebral ischaemia, which is still considerable—it will be extremely difficult to demonstrate an effect of this treatment on outcome. To demonstrate a beneficial effect of such a treatment on outcome at least 2825 patients have to be randomised in a controlled trial. An even larger number of patients would be necessary if treatment is effective on postoperative ischaemia only.

The results of this study do not support the suggestion expressed in reviews that with modern treatment, delayed cerebral ischaemia is the only problem that has to be dealt with to improve outcome in patients with aneurysmal subarachnoid haemorrhage. The explanation is that in two thirds of the patients with poor outcome the causes are the effects of the initial bleeding and rebleeding. Only if these effects can be reduced, will the outcome of all patients improve. Further improvement of the treatment of delayed cerebral and postoperative ischaemia will only have minor effects on the outcome of patients with aneurysmal subarachnoid haemorrhage.

We thank M Mechielsen and A Vet for their excellent data management. We also thank the pharmaceutical firm Pharmacia and Upjohn for their financial support of this study.
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Appendix: list of participating centres (in alphabetical order)

Academic Medical Centre (AMC), Amsterdam (Y Roos, KW Albrecht, M Vermeulen, RJ de Haan)
BovenIJ Ziekenhuis, Amsterdam (A Janssens)
Free University Hospital (VU), Amsterdam (LFM Beenen, FPJN Ploegmakers)
Kennemer Gasthuis, Haarlem (RMCM Janssen, JAM Kuster)
Kennemer Gasthuis, IJmuiden (JA Don, HCW Hoff)
Medisch Centrum Alkmaar (MCA), Alkmaar (MM Vering)
Onze Lieve Vrouwe Gasthuis (OLVG), Amsterdam (P Verlooy)
Rode Kruis Ziekenhuis, Beverwijk (HJMV vd Werd)
Sint Lucas Andreas Ziekenhuis, Amsterdam (JLM Hellenberg Hubar, JAL Vanneste)
Slotervaart Hospital, Amsterdam (RJM Groen)
St Gemini Ziekenhuis, Den Helder (JG Kok)
St Ziekenhuis De Heel, Zaandam (AJ Prazsky, A Koppenaal)
Waterlandziekenhuis, Purmerend (CP Zetsvloot)
Westfries Gasthuis, Hoorn (FJAM Bussemaker, CMB Lambrecht)
Ijsselmeerrakkenhuizen, Lelystad (RJ Tans)
Ziekenhuis Amstelveen, Amstelveen (P Lanting)