Playing the odds in clinical decision making: Lessons from berry aneurysms undetected by magnetic resonance angiography

Article (Published Version)


This version is available from Sussex Research Online: http://sro.sussex.ac.uk/id/eprint/38070/

This document is made available in accordance with publisher policies and may differ from the published version or from the version of record. If you wish to cite this item you are advised to consult the publisher's version. Please see the URL above for details on accessing the published version.

Copyright and reuse:
Sussex Research Online is a digital repository of the research output of the University.

Copyright and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable, the material made available in SRO has been checked for eligibility before being made available.

Copies of full text items generally can be reproduced, displayed or performed and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.

http://sro.sussex.ac.uk
Lesson of the week

Playing the odds in clinical decision making: lessons from berry aneurysms undetected by magnetic resonance angiography

Michael R Johnson, Catriona D Good, William D Penny, Philip RJ Barnes, John W Scadding

The purpose of this report is twofold—to report the potential for magnetic resonance angiography to miss sizeable intracranial aneurysms and to highlight the value of simple, quantitative clinical reasoning when interpreting the results of diagnostic tests.

Subarachnoid haemorrhage accounts for a quarter of all cerebrovascular deaths, and over a third of those who survive have major neurological deficits.6 Intra-cranial aneurysms, the commonest cause of subarachnoid haemorrhage, may present with rupture, mass effect, or, rarely, with emboli phenomena in large aneurysms. The typical presentation of rupture is headache of instantaneous onset that remains continuous and is often associated with nausea, vomiting, meningism, or loss of consciousness. About a third of patients with aneurysmal subarachnoid haemorrhage will re-bleed, and this is a major cause of poor outcome.7,8 The risk of re-bleeding peaks on the first day and then declines.9 Most studies therefore support the need for surgery soon after rupture, and delay in diagnosis or misdiagnosis as migraine or meningitis can have catastrophic consequences.10

Unruptured intracranial aneurysms causing mass effect may present as pain or neurological deficit depending on the site and size of the aneurysm. Such aneurysms are often large or giant,6 and, as most intracranial aneurysms occur at the junction of the internal carotid and posterior communicating artery, the commonest clinical sign is oculomotor palsy. Unruptured intracranial aneurysms causing mass effect are at high risk of subsequent rupture, estimated at 6% a year.11 The optimal method for detecting intracranial aneurysms is intra-arterial digital subtraction angiography. This procedure carries an associated morbidity of transient or permanent neurological disability (of 1% and 0.5% respectively).12 This associated morbidity and increasing access to magnetic resonance imaging has led to interest in the use of magnetic resonance angiography for assessing patients at high risk of symptomatic intracranial aneurysm.

Case reports

Case 1

A 21 year old man developed a headache of instantaneous onset. The headache improved overnight, becoming persistent and centred behind the left eye. Ten days later he noticed drooping of his left eyelid and double vision. Clinical examination revealed a left, complete, oculomotor palsy. Results of magnetic resonance imaging of the head and intracranial magnetic resonance angiography (fig 1) were reported to be normal by a senior consultant neuroradiologist who was aware of the clinical suspicion of a posterior communicating artery aneurysm. The patient's physician reasoned that, since magnetic resonance angiography has a sensitivity to detect intracranial aneurysms of ≥95%, the probability of a posterior communicating artery aneurysm was ≤5%.

The diagnosis of posterior communicating artery aneurysm was discounted, and an alternative cause for the patient's oculomotor palsy was sought. Subsequent failure to identify another cause led to a re-evaluation, and cerebral intra-arterial digital subtraction angiography revealed a left 8 mm posterior communicating aneurysm.9

The probability of a disease following a diagnostic test is critically reliant on prior clinical probability

Division of Neuroscience and Psychological Medicine, Charing Cross Hospital, London W6 8RF
Michael R Johnson consultant neuroradiologist
Wellcome Department of Cognitive Neurology, University College London Institute of Neurology, London WC1N 3BG
Catriona D Good clinical lecturer
William D Penny statistician
National Hospital for Neurology and Neurosurgery, London WC1N 3BG
John W Scadding consultant neuroradiologist
Neurosciences Centre, Kings College Hospital, London SE5 9RS
Philip RJ Barnes consultant neuroradiologist
Correspondence to: M R Johnson m.johnson@ucl.ac.uk
BMJ 1991;302:1547–9

![Fig 1 Case 1: magnetic resonance angiography with maximum intensity projection has normal appearance (top), whereas intra-arterial digital subtraction angiography (injection of left internal carotid artery) shows a large left posterior communicating artery aneurysm (bottom)](https://www.bmj.com/content/322/7290/1347)
Clinical review

aware of the clinical suspicion of intracranial aneurysm and after analysis of maximum intensity projections and axial base images.

The major clinical lesson, however, concerns the degree of reliance placed on a negative test result when clinical features strongly suggested otherwise. In case 1 the patient's initial physician reasoned that, because magnetic resonance angiography has a ≥95% sensitivity to detect intracranial aneurysms, the probability of the patient having an aneurysm was ≤5%. This reasoning fails to consider the probability of intracranial aneurysm based on both the clinical features and the test result. The probability of a diagnosis based on both probabilities is termed the posterior (post-test) probability and is calculated using Bayes's rules.\(^{12}\)

Of key importance in Bayesian calculations is the accurate estimation of prior clinical probability and test sensitivity and specificity. For case 1, we estimate the clinical probability of posterior communicating artery aneurysm to be high (90%). This is based on the fact that the patient's oculomotor palsy involved pupillary fibres (characteristic of a "surgical third") and that isolated painful third nerve palsies are the hallmark of posterior communicating artery aneurysms. The test sensitivity (the ability of magnetic resonance angiography to detect intracranial aneurysms) is ≥95% for aneurysms > 6 mm in diameter or when axial base and spin-echo images are reviewed as well.\(1\) The specificity of magnetic resonance angiography (the probability of a negative result when disease is absent) ranges from 92% to 100%.\(13\) Assuming a prior clinical probability of an aneurysm of 90% and magnetic resonance angiography sensitivity and specificity of 95% and 92% respectively, the probability of a posterior communicating artery aneurysm after a negative magnetic resonance angiography is reduced from 90% to

artery aneurysm (fig 1). The aneurysm was surgically clipped without complication.

Case 2

A 48 year old woman developed a headache of instantaneous onset. The headache improved over the course of a few hours, becoming centred behind the right eye. Two days later she became drowsy, photophobic, and meningitic. Computed tomography of the head gave normal results. Examination of the cerebrospinal fluid revealed bloodstained fluid (opening pressure not recorded), a red blood cell count of 180×10\(^3\)/l, white blood cell count 210×10\(^3\)/l (70% lymphocytes, 30% neutrophils), and glucose concentration 2.2 mmol/l (serum concentration 8.0 mmol/l). A diagnosis of "bloody tap" and meningitis was made, and she received benzylpenicillin.

The next day she deteriorated, and partial oculomotor palsy of the right pupil was noted. The results of magnetic resonance imaging of the head (before and after use of gadolinium contrast agent) and magnetic resonance angiography were reported by a senior consultant neuroradiologist, who was aware of the clinical suspicion of a posterior communicating artery aneurysm, as showing meningeal enhancement consistent with subarachnoid haemorrhage but with no sign of an aneurysm (fig 2). In view of the strong clinical suspicion of intracranial aneurysm, cerebral intra-arterial digital subtraction angiography was performed and revealed a right 8 mm posterior communicating artery aneurysm (fig 2). The aneurysm was surgically clipped without complication.

Comment

These cases highlight the potential for high quality magnetic resonance angiography to miss sizeable intracranial aneurysms and thereby contribute to delayed or wrong diagnosis. In both cases the images were reported by a senior consultant neuroradiologist
32.85% (fig 3). Thus, in these circumstances a negative result from magnetic resonance angiography cannot be used to exclude the diagnosis. The delay in diagnosis occurred because insufficient weight was given to the clinical findings and too much weight to the magnetic resonance angiography result. For case 2, however, once the misdiagnosis of meningitis was discarded, appropriate weight was given to the clinical probability of aneurysm, and intra-arterial digital subtraction angiography was undertaken immediately despite the negative result from magnetic resonance angiography.

We have avoided using the terms positive and negative predictive values. The positive predictive value is the probability a disease is present given a positive test result. The negative predictive value is the probability a disease is absent given a negative result. The probability that a disease is present given a negative result is therefore 1 — negative predictive value. The term posterior probability avoids this potential confusion and simply refers to how likely a patient is to have a disease given the result of a diagnostic test. We could have calculated posterior odds rather than posterior probability, but we have avoided the use of odds and likelihood ratios because they require the conversion of probabilities to odds. As clinical likelihood, sensitivity, and specificity are all usually expressed as probabilities rather than odds, we prefer the more intuitive posterior probability to express the post-test likelihood of a disease.

Fig 4 shows how the probability of a disease after a diagnostic test is critically dependent on the prior clinical probability. For example, a small reduction in the prior clinical probability of an intracranial aneurysm from 90% to 50% reduces the probability of aneurysm given a negative result from magnetic resonance angiography from 33% to 5%. Such a reduction may not be enough to rule out the diagnosis with certainty, but it might be enough to question it. The reliance of posterior probability on prior clinical suspicion is daunting since even small errors in the estimation of clinical suspicion can substantially affect the final decision about whether a disease is present. Methods to quantify clinical suspicion are described elsewhere.13 14 Doctors' diagnostic opinions can differ because some are better than others in their ability to correctly estimate the prior clinical probability of a disease, in their knowledge of test sensitivity and specificity, or in their intuitive ability with Bayesian statistics.

The formulas in fig 3 provide a simple and convenient method for calculating the probability of a disease based on both the prior clinical probability and the result of a diagnostic test.

Contributors: MRJ conceived, researched, and produced the article. CDG contributed data analysis, discussed core ideas, and helped draft and edit the article. WDP contributed statistical expertise, discussed core ideas, and helped draft and edit the article. PKB and JWS supplied clinical details of patients. JWS discussed core ideas and helped draft and edit the article. JWS is guarantor for the article.

Funding: No additional funding.

Competing interests: None declared.

Corrections and clarifications
Minerva
Minerva was perhaps a bit too keen to report a study on doctors committing suicide (17 March). She cited the article as appearing in the Journal of Epidemiology and Community Health, but unfortunately at that point the article had not been published. She failed to realise she was working from prepublication proofs, not a reprint. The paper has now been published—in volume 55, pp 296-300. Thanks to the reader who alerted us to this error.

ABC of hypertension: Blood pressure measurement: Part 1—Sphygmomanometry: factors common to all techniques
The caption to the diagram on p 981 of this article by Gareth Beevers and colleagues (21 April, pp 981-5) wrongly described the blood pressure pattern as normal. It should have read: “Example of ambulatory blood pressure pattern plotted by the DABL® Program showing a marked variability of blood pressure.”


(Accepted 17 January 2001)