MR venography in idiopathic intracranial hypertension: unappreciated and misunderstood

J N P Higgins, J H Gillard, B K Owler, K Harkness, J D Pickard

Background: Venous sinus disease must be excluded before diagnosing idiopathic intracranial hypertension but is found only rarely in typical cases. Magnetic resonance venography (MRV) is the technique of choice for investigating this, and provides images that are diagnostic and easy to interpret. However, recent work using more invasive techniques has documented pressure gradients and stenoses in the lateral venous sinuses in many cases of idiopathic intracranial hypertension.

Objective: To examine the reason for this discrepancy and to establish whether there are characteristic appearances on MRV in idiopathic intracranial hypertension that are routinely overlooked in clinical practice.

Methods: MRV’s from 20 patients with idiopathic intracranial hypertension were reviewed, unblinded, by two neuroradiologists, and their appearances rated for focal narrowings and signal gaps. A control group of 40 asymptomatic volunteers, matched for age and sex with the patient group, was recruited prospectively for MRV, and their scans rated in the same way.

Results: The lateral sinuses presented a range of appearances with quite different distributions in the two groups (p<0.001). Bilateral lateral sinus flow gaps were seen in 13 of 20 patients with idiopathic intracranial hypertension and in none of 40 controls.

Conclusions: A historical failure to use normal healthy controls to establish the boundaries between imaging artefact, normal anatomical variant, and disease means that the pathological significance of the different appearances of the lateral sinuses on MRV has not so far been appreciated.

This study was carried out with a view to establishing whether there are characteristic appearances on MRV in idiopathic intracranial hypertension that are being routinely overlooked in clinical practice, and so why.

METHODS

Patient group
All patients coded with a diagnosis of idiopathic intracranial hypertension between January 1998 and January 2001 at Addenbrooke’s Hospital and the Norfolk and Norwich Hospital were identified. Their notes were retrieved by the audit departments at both hospitals and manually searched by two of us (BKO and KH). Patients were diagnosed as having idiopathic intracranial hypertension if they had presented with a syndrome of raised intracranial pressure, usually manifest by headache and/or visual disturbance, without ventricular enlargement or an intracranial mass on imaging, with no evidence of venous sinus thrombosis, and with normal cerebrospinal fluid (CSF) constituents. Only patients who had undergone MRI and MRV as part of their diagnostic work up were included in the study.

Twenty two cases were identified. One patient whose diagnosis was subsequently changed to tension headache was excluded. Another, in whom the CSF pressure had never been measured, was also excluded. This left 20 patients who had had MRI and MRV, all of whom had been reported as showing no evidence of venous sinus disease (table 1). These scans were reviewed by two neuroradiologists together (JNP and JHG) to look for evidence of acute thrombosis on the static images and abnormalities of flow on MRV. The presence of thrombosis was judged by standard diagnostic

Abbreviations: MRV, magnetic resonance venography
The MRV appearances of the superior sagittal sinus and each lateral sinus were scored for each patient according to whether they were normal (0), had one or more areas of focal narrowing (1), or contained one or more signal gaps (2). A superior sagittal sinus that was uniformly narrowed over its anterior third was regarded as normal. A lateral sinus that appeared uniformly narrow was also regarded as normal. Superior sagittal sinus scores were considered separately from lateral sinus scores. Lateral sinus scores were summated in each individual to give values from 0 (normal) to 4 (bilateral signal gaps) (fig 1).

Control group
Normal controls were recruited over a period of 18 months by poster advertisement or from the patient waiting list for non-cerebral MRI and had none of the standard contraindications for MRI. All had MRI and MRV but first were screened by one of us (JNPH or JHG) using a questionnaire for symptoms relevant to their status as controls and categorised either as true (asymptomatic) normal or false (symptomatic) normal. Volunteers were informed before screening that their answers would not affect their eligibility for the study or a scan. Recruitment continued until there were 40 true normal subjects, matched for sex and for age within three years, with the patient group (table 2). The MRIs and MRVs of this group were reviewed subsequently by two neuroradiologists (JNPH and JHG) and scored in the same way as the patient group.

MRI and MRV
All patients had had dual echo and T1 weighted MRI of the brain and either phase contrast or time of flight MRV. In-house scans were done on a GE 1.5 T Signa magnet using three dimensional phase contrast MRV, usually with velocity encoding of 15 cm/s but occasionally using 30 cm/s and once 40 cm/s. Separate acquisitions were made for the sagittal and lateral sinuses. Scans undertaken at the referring hospitals on a variety of machines were not repeated if they were of good quality.

All volunteers were scanned in-house on the same GE 1.5 T Signa magnet. All had sagittal T1 and axial T2 MRI. MRV was done using a three dimensional phase contrast technique with 15 cm/s velocity encoding (time of repetition 25 ms; flip angle 20°; time of echo 7.2 ms). A sagittal slab of 12 contiguous slices, each 2.2 mm thick, was used to image the superior sagittal sinus and other midline venous structures. An axial slab of 28 slices, each 2.2 mm thick, was used to image the lateral sinuses. Scans were displayed as maximum intensity projection reconstructions for rating, with the source images available to resolve any uncertainty created by overlapping vessels.

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<tr>
<th>Table 1 Demographic details of 20 patients with idiopathic intracranial hypertension</th>
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<tr>
<td>17 female, age 16 to 69 years (mean 30)</td>
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<td>3 male, age 17 to 57 years (mean 42)</td>
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<td>Body mass indices: 21–47 kg/m² (mean 35)</td>
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<td>Duration of symptoms before MRI: 1 week to 4 years (mean 5 months)</td>
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<td>CSF pressures (cm H₂O): 23 to 58 (mean 35)</td>
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<th>Table 2 Demographic details of 40 asymptomatic normal volunteers</th>
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<td>34 female, age 14 to 71 years (mean 30)</td>
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<td>6 male, age 15 to 60 years (mean 42)</td>
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Figure 1  Lateral sinus scores and magnetic resonance venography (MRV) appearances. Three dimensional phase contrast MRV; axial views of lateral sinuses. Each lateral sinus scored either normal (0), with one or more focal narrowings (1), or with one or more signal gaps (2) and the results summated. (A) Score 0. (B) Score 1: focal narrowing right transverse sinus, arrow. (C) Score 2: focal narrowing both transverse sinuses, arrows. (D) Score 2: normal right transverse sinus, signal gap left transverse sinus, arrows. (E) Score 3: signal gap right transverse sinus—confirmed on source images, arrow; focal narrowing left transverse sinus, arrow. (F) Score 4: bilateral transverse sinus signal gaps, arrows. SS, right sigmoid sinus; TS, right transverse sinus.
RESULTS
One hundred and forty volunteers were imaged, of whom 56 were designated as true normal; 46 of those could be matched with the patient group. The first 40 were recruited into the study. True normal controls only exceptionally complained of headache. Those excluded could not be age matched, or gave a history of headaches or other illness symptoms.

In the patient group there was no evidence of sinus thrombosis on the static images. On MRV the superior sagittal sinus showed uniform signal throughout its length in 18 cases and appeared relatively hypoplastic over its anterior third in two. The lateral sinuses showed a variety of appearances, the component being flow gaps present on both sides (score 4, fig 1F). Only one patient had strong uniform signal throughout both lateral sinuses (score 0, fig 1A). Figure 2 shows the frequency of the different appearances of the lateral sinuses in this group expressed in their scores as defined above.

In the control group static imaging was normal. On MRV the sagittal sinus showed an area of focal narrowing in five cases. There was some overlap with the patient group in the appearances of the lateral sinuses, but the distribution was significantly different (p<0.001 using the χ² test for independence). The large majority of controls had strong uniform signal from at least one lateral sinus, and usually from both (score 0 or 1). In no control cases were flow gaps seen on both sides (fig 3).

DISCUSSION
In a patient with the appropriate clinical phenotype, the diagnosis of idiopathic intracranial hypertension is essentially one of exclusion of known causes of raised intracranial pressure. This has in many respects become easy with modern imaging. MRI will detect an intracranial mass or MRV. Chronic thrombosis, or partially recanalised thrombus—is the accuracy with which the radiological imaging is indistinguishable from idiopathic intracranial hypertension—is the accuracy with which the radiological imaging excludes venous sinus disease. Once considered, acute thrombosis is not usually difficult to diagnose on the static images and will be confirmed by abnormalities of flow on MRV. Chronic thrombosis, or partially recanalised thrombus, is less obvious on the static MRV but may be recognisable on MRV. In this respect, MRV of the superior sagittal sinus does not normally present any problems in interpretation, a normal study generally giving uniform signal over the course of the sinus, thrombosis—new or old—usually producing quite extensive areas of reduced or absent signal. The lateral sinuses, however, are more difficult to assess, seeming to present a wide variation in the normal anatomy. Either side may be dominant, either may be absent, and frequently they contain filling defects. Ayanzen et al carried out MRV on 100 patients with normal MRI. They found defects in the lateral sinuses in 31%, which they cautioned should not be mistaken for dural sinus thrombosis. Lee and Brazis, making allowances for similar appearances, found no evidence of sinus thrombosis in 20 patients with idiopathic intracranial hypertension.

These studies reflect current practice in the interpretation of MRV but make significant assumptions, the most important being that for the purpose of studying MRV a normal MRI qualifies a patient as a normal control. This is highly questionable. Ayanzen’s patients were undiagnosed and their symptoms unspecified but it is probable—given that their MRI scans were normal—that many were being investigated for headache. Their conclusion ought therefore to have been that lateral sinus defects may be seen in 31% of patients with otherwise normal imaging, many of whom will probably have been referred for headache. Rao and Higgins examined the symptoms of 100 consecutive patients whose MRI scans were normal. They found that 31% complained of headache and 22% of focal neurological symptoms. Seven per cent had acute seizures. These patients cannot be representative of the normal healthy population and particularly not of healthy venous sinuses. Headache, for example, is the cardinal symptom of venous sinus thrombosis and, moreover, has often been present for months before the diagnosis is made.

There must therefore be a suspicion that our understanding of the normal appearances of the venous sinuses, based in part on clinical experience but mirrored in published reports, may be compromised by contamination of control groups with patients who have unrecognised venous sinus disease. Could this explain the failure of MRV to identify abnormalities of the venous sinuses in idiopathic intracranial hypertension?

To address this concern we selected a control group of “supernormals” from our volunteers. These subjects, whom we called true normal, only exceptionally ever suffered from headache and complained of no consistent symptoms that could conceivably be referred back to the head. Headache is a common symptom and inevitably there was a degree of subjectivity in assignment to one or other group. In the interests of the study objectives, however, admission to the true normal group was difficult, only one third of volunteers being considered sufficiently asymptomatic for inclusion in this category. Clearly, many truly healthy subjects must have been rejected from the true normal group (assuming that the symptoms they expressed were not indicative of pathology). These subjects would probably dilute any MRV abnormalities to be found in any analysis of the false normal group. However, their exclusion from the true normal group would have had no significant effect on the composition of this
group, only on the time and effort required to attain the number of subjects needed for the study. On the other hand, also excluded from this group was a substantial proportion of volunteers who complained of quite significant symptoms, sometimes including severe and frequent headache. Some of these had attended for reassurance. Some had been investigated for the symptoms they described but given no specific diagnosis. One subject (who had previously had normal brain CT) had attended the ophthalmology clinic with a diagnosis of possible idiopathic intracranial hypertension.

The result of this selection process has been to expose clear differences between our two sample populations. We found that the great majority of patients with idiopathic intracranial hypertension presented a distinctive pattern on MRV—signal gaps in both lateral sinuses—which was not seen in the control group. A smaller number had lesser defects but these were still unusual in the control group. Moreover, our control group showed substantially fewer defects in the lateral sinuses on MRV than recorded previously, lending credence to the concerns expressed above and suggesting that a re-evaluation of normative data has become necessary.

What do these signal gaps mean? The signal in phase contrast MRV mainly comes from the bulk movement of protons (in blood) within a range of velocities chosen by the operator. As such, it is more a demonstration of flow than of anatomy, although some anatomical information is inevitably present in any study. A segment of signal void in a sinus implies that flow velocities over that segment are outside the range prescribed in the study. This does not necessarily mean thrombosis, and it may not necessarily be abnormal—arachnoid granulations might cause a local alteration of blood flow around them, for example. However, it does raise the possibility of stenosis or occlusion.

In this context King et al and Karahalios et al, using more invasive techniques, have investigated the possibility that unrecognised venous sinus disease may be a frequent cause of idiopathic intracranial hypertension. With angiographic catheters passed into the intracranial venous sinuses through the jugular veins these workers found raised intrasinus pressures in many patients with idiopathic intracranial hypertension. Sometimes these high pressures appeared to be secondary to central venous hypertension but more often they seemed to be the result of stenotic lesions in the lateral sinuses obstructing cerebral venous outflow. This has led one group to propose intracranial venous hypertension as the final common pathway in the aetiology of idiopathic intracranial hypertension. Others, however, have since suggested that the sinus abnormalities observed in these cases are not the cause but the results of raised intracranial pressure (of unknown aetiology) inducing secondary sinus collapse.

Clearly, this latter issue is fundamental to questions regarding the aetiology of idiopathic intracranial hypertension and is not resolved by the current study. However, at least it is now possible to reconcile the non-invasive imaging with catheter venography and manometry. This means that, regardless of cause or effect, patients with idiopathic intracranial hypertension who have bilateral lateral sinus defects on MRV are likely to have raised venous pressures and stenotic lesions in their lateral sinuses. It means that, even in patients in whom frank venous sinus thrombosis has been excluded, MRV may still present an appearance that equates with raised intracranial pressure (an idea already explored by Quattrone et al with respect to chronic daily headache). The possibility that this appearance may sometimes be indicating the cause of raised intracranial pressure is suggested by recent work documenting clinical improvement in cases of idiopathic intracranial hypertension following dilatation and stenting of the lateral sinuses. This pattern on MRV may thus in the future direct further investigation of these patients, and this concept should encourage a refinement of imaging techniques.

Our patients were identified blind to a review of their MRI and MRV (although in the knowledge that they must have been reported as showing no evidence of sinus thrombosis). True normal controls were identified by interview before they were scanned. Otherwise this study was unblinded. Patients had a variety of non-standardised MRI and MRV examinations. Controls were scanned according to a defined protocol. It would have been impossible to blind the observers to the imaging of patients versus controls. Inevitably, therefore, there will have been some bias in the reporting of the two groups, but the differences between them are so striking that this is unlikely to have influenced the results significantly.

Conclusions

MRV in patients with idiopathic intracranial hypertension commonly presents a pattern of bilateral lateral sinus defects rarely seen in asymptomatic controls. This result, at variance with published reports, implies a historical failure to separate patients with unrecognised venous sinus disease from healthy subjects in control populations, and implies a widespread failure to differentiate between normal variant and disease on MRV in clinical practice. This is highly relevant in the light of recent proposals regarding the aetiology and management of this condition.

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REFERENCES


