MATTERS ARISING

MRI in monitoring the treatment of multiple sclerosis: concerted action guidelines

We would like to comment on the article by DH Miller et al. At the University of British Columbia, (UBC) we have been involved in serial MRI evaluations of multiple sclerosis (MS) since 1985. We originally did our serial MRI evaluations without the use of gadolinium and at present we are using gadolinium in selected cases. Dr Miller and his colleagues at Queen Square have done elegant studies both with and without gadolinium to detect activity of MS lesions by using MRI.

There is no doubt that gadolinium can show breakdown of the blood brain barrier (BBB) in new and enlarging MS lesions. Gadolinium may even show BBB impairment in some old lesions in which no changes can be seen on the standard unenhanced scans. In Dr Miller’s experience this occurred in 10% of the instances. Conversely, the periventricular lesions seen on standard unenhanced scans, but missed on gadolinium enhanced scans, is higher. Dr Miller et al have shown that in patients with a secondary progressive and early relapsing remitting MS, about 20% of new lesions did not enhance.

In patients with benign MS and primary progressive MS, this percentage is even higher (67% and 95% respectively). They have also shown that in 25% of enhancing lesions the enhancement persisted for less than 4 weeks. Therefore one might miss detecting activity in up to 25% of lesions with a 4 weekly scanning interval. We have found using a low field strength unit at UBC (0.15 Tesla), and without gadolinium, that only 67% of active lesions were identified if the scanning interval is 4 weeks rather than 2 weeks. If the scanning interval is 6 weeks, only 40% of the active lesion is identified.

Our experience therefore suggests to us that to maximise detection of MS lesions on MRI, scanning at shorter intervals will allow for a higher detection rate. The savings in time and cost incurred by not doing gadolinium enhanced scans could then be applied to scanning the patient more frequently, obviously within the limits of patient tolerance. A single set of multi-echo T2 weighted transverse images carefully repositioned is all that is required to monitor the morphological changes that we see. To perform gadolinium studies, two additional T1 weighted spin echo sequences are required. The imaging time is therefore increased at least two fold.

The addition of gadolinium on a routine basis also increases the invasiveness of the procedure. Not only must material that is potentially harmful be injected (there are rare reports of anaphylaxis) but patients also report discomfort from the injection. Our patients have complained of headaches and other transient symptoms related to the injection of gadolinium. Fortunately we have not had any serious side effects. Nevertheless, patients are probably more willing to be scanned at more frequent intervals if an injection is not required.

Those of us at UBC who have been doing serial MRI evaluations for many years feel therefore that for routine clinical trial evaluation purposes, frequent unenhanced scans will suffice. However, if one is doing a study in which evaluation of the immediate therapeutic effect on leakage of BBB is an important endpoint, then gadolinium must, of course, be used.

DH PATY
Division of Neurology

Miller et al reply:
We welcome the comments of Professor Paty and his colleagues, whose MRI studies have contributed much to our understanding of the natural history of MS and of the use of MRI in diagnosing and monitoring the disease. We agree with Paty that frequent serial unenhanced T2-weighted brain MRI is a powerful tool in monitoring MS treatment trials, and we also recommend that it is performed on every occasion.

Another limitation is the low pathological specificity of MRI. On unenhanced scans, oedema, inflammation and gliosis all produce similar signal changes. An added benefit of gadolinium enhancement is that it identifies the active, inflammatory MS lesion, and may give some indication of the mode of action of the drug under study. The risk of anaphylaxis from gadolinium-DTPA, although remote (about 1/100 000), should always be discussed with patients when obtaining informed consent. In our experience, the minor side effects referred to by Paty have been both rare and nontroublesome.

Finally, we agree that patient compliance is essential to a successful study, and it might be argued that a 40 minute monthly scan is more acceptable than a 20 minute fortnightly scan.

DH MILLER
P BARKHOF
I BERRY
L KAPPOS
G SCOTTI
AJ THOMPSON
University Department of Clinical Neurology, Institute of Neurology


The prognosis of primary intracerebral tumour presenting with epilepsy: the outcome of medical and surgical management.

We welcome the paper by Smith et al which attempts to address the importance of resective surgery in intrinsic brain tumours presenting with epilepsy as the first symptom. We agree that it is important to consider the timing of surgery in this group of patients, but definitions of “early” and “late” in this context have not yet been agreed. There are, however, a number of methodological problems posed in this paper, which we do not allow the correct comparisons to be made.

First, a definition of “resective surgery” is not provided, consequently, no distinction is made between “total” versus “subtotal” resection. This comparison is a major prognostic factor reported by Laws et al. Thus, it is not the extent of surgery which is important as suggested in the author’s discussion, but whether the tumour is considered to be totally removed or not. Unfortunately, the authors fail to make this important distinction.

Second, no direct comparison of duration of survival is made between “resective” surgery versus “non-resective” surgery in the first symptom epilepsy group, who demonstrated a non-enhancing CT scan lesion associated with a low grade tumour at the time of surgery. This is an important comparison, because it is likely that patients presenting with epilepsy who are found to have non-enhancing lesions on CT scan will have low grade tumours. Depending on their site in the brain, some non-enhancing CT, low grade tumours are likely to be totally resectable. Previous studies of low grade tumours have demonstrated that non-enhancing CT lesions and total resection both carry a better prognosis.

Presumably these important comparisons were not made, because the number of first symptom epilepsy group undergoing any form of surgery was 65, and only 20 had any type of resection. Subdivision into patient’s suffering from low grade tumours demonstrated as a non-enhancing lesion on CT scan, who had surgery compared with those that did not; and those that had “total” versus “subtotal” resection with this type of lesion, would probably have resulted in numbers too small for statistical comparisons to be made.

This paper illustrates the problem of trying to use data from a single centre in order to make definitive management policies. It emphasises the importance of considering collecting a national data base to pose the important questions raised by the authors of this paper.

JOHN N WILDE
Department of Neurosurgery, Alexandra Wing, The Royal London Hospital, London E1 1BB, UK

1 Laws ER, Tawel WF, Clifton MB, Ozakazi H. 61. Neurosurgical management of low grade