Pre-operative verbal memory fMRI predicts post-operative memory decline after left temporal lobe resection

Mark P. Richardson,¹ Bryan A. Strange,² Pamela J. Thompson,³ Sallie A. Baxendale,³ John S. Duncan^{1,3} and Raymond J. Dolan²

Correspondence to: Dr Mark Richardson, Box 29, Institute of Neurology, Queen Square, London WC1N 3BG UK E-mail: m.richardson@ion.ucl.ac.uk

Summary

Functional MRI (fMRI) of cognitive tasks depends on technology widely available in the clinical sphere, but has yet to show a role in the investigation of patients. We report here the first demonstration of a clinically valuable role for cognitive fMRI. Temporal lobe epilepsy (TLE) is commonly caused by hippocampal sclerosis and is frequently resistant to drug treatment. Surgical resection of the left hippocampus in this setting can cure seizures, but may produce significant verbal memory decline, which is hard to predict. We report

10 right-handed TLE patients with left hippocampal sclerosis who underwent left hippocampal resection. We compared currently used data for the prediction of post-operative verbal memory decline in such patients with a novel fMRI assessment of verbal memory encoding. Multiple regression analyses showed that fMRI provided the strongest independent predictor of memory outcome after surgery. At the individual subject level, the fMRI data had high positive predictive value for memory decline.

Keywords: fMRI; memory; hippocampus; epilepsy; surgery

Abbreviations: fMRI = functional MRI; IAP = intracarotid amytal procedure; MTLE = mesial temporal lobe epilepsy *Received May 6, 2004. Revised July 9, 2004. Accepted July 10, 2004. Advanced Access publication September 30, 2004*

Introduction

Epilepsy is the most prevalent serious neurological disorder, with a prevalence of 0.5–1% (Hauser, 1998). Approximately 30% of people with epilepsy continue to have seizures despite antiepileptic drug treatment; the majority suffer from focal epilepsy (Crawford, 2000). In this context, neurosurgery to resect the epileptic focus is an important and under-used treatment option. The first randomized trial of surgery versus optimal drug treatment in temporal lobe epilepsy (Wiebe *et al.*, 2001) demonstrated a seven-fold increase in the likelihood of being seizure-free in the surgically treated group, with an associated highly significant improvement in quality of life.

The most frequent pathology identified in drug resistant epilepsy patients is hippocampal sclerosis, giving rise to the common syndrome of mesial temporal lobe epilepsy (MTLE). Despite a seizure-free outcome, many patients suffer a significant decline in memory ability after temporal lobe resection, especially for verbal memory following left-sided

resection in right-handed patients. Decline correlates inversely with severity of hippocampal sclerosis in the surgical resection specimen (Hermann *et al.*, 1992; Sass *et al.*, 1994) and with severity of hippocampal sclerosis assessed by left hippocampal volume measured on pre-operative MRI (Trenerry *et al.*, 1993).

Pre-operative memory performance is also a predictor of post-operative memory decline, with better performance predicting worse decline (Helmstaedter and Elger, 1996; Jokeit *et al.*, 1997). Concern about memory decline and current limitations in its prediction play an important role in limiting the number of patients undergoing potentially seizure-curing surgery. Memory may suffer in this context because hippocampus and related mesial temporal lobe (MTL) structures are crucial to long-term episodic memory function (Squire and Zola-Morgan, 1991; Squire, 1992; Lepage *et al.*, 1998; Cabeza and Nyberg, 2000).

¹Department of Clinical and Experimental Epilepsy and ²Wellcome Department of Imaging Neuroscience, Institute of Neurology, University College London, and ³National Society for Epilepsy Chalfont Centre, Chalfont St Peter, UK

Patients judged at risk of post-operative amnesia undergo the intracarotid amytal procedure (IAP or 'Wada test'), during which sodium amytal is injected into each internal carotid artery in turn, allowing selective anaesthesia of each cerebral hemisphere. The ensuing period of unilateral hemispheric anaesthesia enables the memory capacity of the un-anaesthetized hemisphere to be examined in isolation. This test is crude and limited due to of the brevity of the anaesthesia, dysphasia during dominant hemisphere injection and patient sedation. It is also highly invasive and carries a significant risk of adverse outcomes, including stroke. Although IAP provides a reasonable index of the risk for amnesia, it is a poor predictor of the extent of verbal memory decline, although some studies have shown modest predictive value (Loring *et al.*, 1995; Jokeit *et al.*, 1997; Lee *et al.*, 2003).

Functional MRI (fMRI) is attractive as a tool to examine memory in patients because of its wide availability and minimal risks. A small number of studies has shown patients with left MTLE preferentially activate right hippocampus in tasks assumed to require left or bilateral hippocampal function (Detre *et al.*, 1998; Dupont *et al.*, 2000; Jokeit *et al.*, 2001; Golby *et al.*, 2002). None of these studies used a design that allowed memory specifically to be examined in isolation.

Only one previous study from our group (Richardson *et al.*, 2003) has shown that right-handed patients with left hippocampal sclerosis preferentially activate the right hippocampal formation in a verbal memory encoding task. We also showed that, in left hippocampal sclerosis patients, there is a distribution of activity between left and right hippocampi during this verbal encoding task that reflects the severity of left hippocampal sclerosis. Thus, patients with mildly affected left hippocampi showed relatively greater activity in left hippocampus than right hippocampus while patients with more severe pathology showed relatively more activity in right than left hippocampus (Richardson *et al.*, 2004).

Two studies have revealed the potential for fMRI of a memory-related task to predict clinical outcomes in MTLE patients: (i) fMRI combined with IAP correctly predicted which patients undergoing anterior temporal lobe resection would become seizure-free (Killgore *et al.*, 1999); and (ii) in the same group of patients, there was a correlation between mesial temporal fMRI activity and recognition memory change following surgery (Casasanto *et al.*, 2001).

To date, there are no clear demonstrations that fMRI has a better predictive value for clinical outcomes compared with other established tests in any neurological field. Our aim in this study was to investigate the utility of event-related fMRI as a predictor of verbal memory decline in right-handed patients with left hippocampal sclerosis undergoing left anterior temporal lobe resection and to compare fMRI with existing techniques.

We previously recruited 26 right-handed patients with left hippocampal sclerosis who underwent evaluation for possible surgery in our centre (the pre-operative data for 24 of these were reported previously (Richardson *et al.*, 2003, 2004). Ten of these subjects underwent surgery, became seizure-free and

had histopathologically proven hippocampal sclerosis. The predictive value of pre-operative fMRI for post-operative memory decline in these subjects is reported here.

In this study, we examined encoding of neutral words. We show that the severity of post-operative verbal memory decline is strongly predicted by a multiple regression model which includes left hippocampal volume, pre-operative verbal memory score and the difference in successful encoding activity between left and right hippocampus. We show that hippocampal encoding activity difference is the strongest independent predictor and has high predictive value at the level of individual patients.

Methods Subjects

Twenty-six consecutive subjects with an MRI-based diagnosis of left hippocampal sclerosis were recruited prospectively from the epilepsy surgery programme of the National Hospital for Neurology and Neurosurgery, London, UK. All 26 subjects underwent an fMRI study during evaluation for surgery.

Inclusion criteria for the 10 subjects reported here were: drug treatment resistant MTLE; right-handed; normal right hippocampal imaging parameters; first language English; seizure-free at least 6 months following left temporal lobe resection; and histopathology of the resection specimen showed hippocampal sclerosis. We included all the subjects who fulfilled these criteria. In the other 16 subjects, surgery was deferred because of a number of factors, which included: significant risk of substantial memory decline or amnesia (according to structural imaging, neuropsychometry and IAP); patient choice; and substantial improvement in seizure control following a change in antiepileptic drug treatment. Participants gave written consent and the study was approved by the Joint Research Ethics Committee of the Institute of Neurology and National Hospital for Neurology and Neurosurgery.

Pre-operative neuropsychometry

To assess verbal memory, we used the List Learning and Story Recall subtests of the Adult Memory and Information Processing Battery (Coughlan and Hollows, 1985) as part of a larger neuropsychometric test battery used for routine pre-surgical evaluation. The List Learning test has an initial registration component (maximum score = 75) and a delayed component (maximum score = 15). The Story Recall Test compares immediate and delayed recall of a story and yields a percent retained score (maximum score = 100). We obtained three clinical neuropsychological measures: (i) list learning—immediate (pre-operative immediate list recall); (ii) list learning—delayed (pre-operative delayed list recall); and (iii) story recall (pre-operative story recall).

Pre-operative structural imaging

Structural MRI was carried out at 1.5 T (Horizon Echospeed, General Electric, Wilwaukee, WI, USA) as part of routine pre-surgical evaluation, including T1 volume and dual-echo whole brain T2-map (Duncan *et al.*, 1996). Hippocampal volumetry was carried out according to a previously published protocol (Van Paesschen *et al.*, 1995).

Pre-operative functional imaging

Functional imaging was performed according to a previously described method (Richardson *et al.*, 2003, 2004). In brief, subjects were scanned at 2 T (Siemens VISION, Siemens, Erlangen, Germany), acquiring T2*-weighted image echo planar imaging (EPI) volumes, providing blood-oxygenation-dependent (BOLD) contrast [33 slices; whole brain; voxel dimensions = $3 \times 3 \times 3.67$ mm; TE (echo time) = 40 ms; TR (repetition time) =2.5 s]. SPM99 was used for image analysis (Friston *et al.*, 1995). The images were realigned, corrected for slice timing differences, transformed to the standard anatomical volume and smoothed (8 mm kernel).

A verbal encoding task was used. During scanning subjects were visually presented with 255 single words, including 36 emotionally aversive words (e.g. 'cancer', 'rape', 'terrorist') (Strange *et al.*, 2000), one every 4.5 s. The word pool from which these were drawn is available on request. Subjects pressed a right-hand button to indicate whether the word was 'living' or 'non-living', and were not asked to memorize the words. Ninety minutes after scanning, subjects performed a surprise recognition memory test (not scanned): subjects were asked to indicate whether the word was definitely remembered (R response); if the word seemed familiar (K response); or was new (N response) (Tulving, 1985). The encoding stimuli were then conditionalized according to subjects' recognition responses. Recognition accuracy (D') was calculated for stimuli labelled 'R' as (hit rate) – (false alarm rate).

To test for subsequent memory effects, imaging data were analysed within a two-level random-effects analysis employing an event-related design (Friston *et al.*, 1998). At the first level, trial-specific responses were modelled and each subject's movement parameters were included as confounds. Contrasts of parameter estimates were calculated to produce a 'contrast image' for each subject of R minus K for neutral items only; we have showed previously that the responses to emotional words were dependent on amygdala pathology, which was not correlated with hippocampal pathology (Richardson *et al.*, 2004).

We created for each patient a voxel-by-voxel image of left minus right encoding activity (R minus K) difference; these 'encoding asymmetry' images were used for further analyses. At the second level, simple regression was used to examine effects within the group. We chose P < 0.05 corrected for peak height both across the whole brain and within the small volume of left hippocampus using a 5 mm radius sphere centred on the peak activation in the left hippocampus in normal subjects in our previous study (Richardson et al., 2003) as the threshold for significance.

Post-operative neuropsychometry

At 3 months post surgery, each subject was tested on parallel versions of the List Learning and Story Recall; these parallel versions have standardized equivalent difficulty. We again collected three neuropsychological measures: (i) list learning—immediate (post-operative immediate list recall); (ii) list learning—delayed (post-operative delayed list recall); and (iii) story recall (post-operative story recall). Using these three measures, performed pre-operatively and repeated post-operatively, we calculated measures of verbal memory change between the first pre-operative and second post-operative assessment (pre-operative—post-operative change in immediate list recall, pre-operative—post-operative change in delayed list recall, pre-operative—post-operative change in story recall).

Data reduction

We expected that there would be a high degree of intercorrelation amongst the measures of memory. Therefore, we used principal components analysis to identify a factor accounting for the largest component of variance amongst these scores. Thus, the three pre-operative measures (pre-operative immediate list recall, pre-operative delayed list recall, pre-operative story recall) were entered into a principal components analysis, from which the first principal component was extracted. Likewise, the three memory change measures (pre-operative—post-operative change in immediate list recall, pre-operative—post-operative change in delayed list recall, pre-operative—post-operative change in story recall) were entered into a principal components analysis, from which the first principal component was extracted.

Prediction of change in memory score: model optimization

We entered the variables predictive of verbal memory outcome into a stepwise linear regression to identify the most important predictive variables.

Hypotheses to be tested

We tested the following hypotheses:

- Pre-operative verbal memory predicts verbal memory decline after surgery.
- (ii) Left hippocampal volume predicts verbal memory decline after surgery.
- (iii) Encoding asymmetry (derived from fMRI data) predicts verbal memory decline after surgery.
- (iv) Encoding asymmetry (derived from fMRI data) is the best predictor of verbal memory decline after surgery.

Results

Demographic, clinical and memory data are summarized in Table 1. Nine of the 10 patients showed decline in at least two of the three memory measures between pre-operative scores and post-operative scores. There was decline in immediate list recall in six patients, no change in two and a slight improvement in two (mean change between pre-operative and postoperative scores was a decline of 5.7 points, ranging from a decline of 17 points to an improvement of 3 points; P = 0.028, paired t-test, 2-tailed). The delayed list recall showed decline in all but one, who had no change (mean change between preoperative and post-operative scores was a decline of 3.6 points, ranging from a decline of 9 points to zero decline, P = 0.002, paired t-test, 2-tailed). There was decline in story recall in six patients and improvement in four, although overall this change was not significant across the group (mean change between pre-operative and post-operative scores was a decline of 14.2 points, ranging from a decline of 72 points to an improvement of 93 points, P = 0.4 paired t-test, 2-tailed).

Co-variation between memory scores

The recognition memory score derived from the recognition test carried out following pre-operative fMRI (D') correlated

Table 1 Demographic and clinical data for the subjects studied

Pre- Pre- Post- Post- Post- Recognition operative operative operative accuracy delayed story immediate delayed story following list recall list recall encoding recall recall fMRI	0.416	0.518	0.324	0.373	0.440	0.603	0.427	0.318	0.303	0.092
Post- e operativ story recall	100	100	14	40	133	75	50	71	14	37
Post- operative delayed list recall	5	∞	0	9) vo	∞	2	7	8	7
Post- Post- operative operativ immediate delayed list recall list recall	26	47	36	49	, 4	50	39	40	27	24
Pre- toperative story recall	83	69	65	92	. 04	94	91	92	98	54
	6	==	S	10	9	∞	11	11	∞	ω
Pre- operative immediate list recall	39	59	36	49	. 4	84	56	45	35	21
Age at Pre- Pre- completion operative operativ of full-time immediate delayed education list recall list (years) recall	81	91	20	9	91	81	61	81	16	16
VIQ PIQ Age at Pre- (WAIS-R) (WAIS-R) completion operative of full-time immediate education list recall (years)	112	73	110	19			117	102	92	68
VIQ I	86	71	102	73	82	68	90	92	92	83
Drugs V and (dose (mg per day)	CBZ400	VPA1500	0 8		0	TOP325 CBZ1800	CLB10 CBZ800	VPA1200 LTG400	TOP400 GBP3600	LTG200 CLB10 CBZ2000 CLB30
Seizure types (frequency per month)	mTL	CFS (4) mTL CPs (3)	CFS (3) mTL CPS (8)	(S) ILW	CPS (1)	CPS (20) mTL	CPS (5) mTL	CPS (2) mTL	CPS (12) mTL	CPS (1), SG TCS (1) mTL CPS (0.5)
	0.4	0.75	26	'n	, 15	∞	9	4	4.5	2.5
Age at Age at first febrile onset of convulsion epilepsy (months) (years)	5	6	11		15	12	15	12		30
Patient Sex LHV LHT2 Age Aetiology (mm ³) (ms)	Febrile	Febrile	Febrile Convulsion	E	Febrile	convulsion Febrile	convulsion Febrile	convulsion Febrile	convulsion None known	Febrile convulsion
Age A	30 F	23 F	4 2 F 2	74 Z	. 45 F		CC 28 F	20 F	28 N	38 FF
.HT2 .	96.4	95.7	94.1	92.4	86.5		94.8	96.4	94.8	
LHV LHT (mm³) (ms)	2026 9	1819 9	1408 9	1890 9			2366 9	1600	1762 9	1779 93.9
Sex 1	F 2	M	M	TT -		Т	M 2	Т	M	Т
Patient	1	6	8	4	· v	9	7	∞	6	10

CLB = clobazam; CBZ = carbamazepine; GBP = gabapentin; LHT2 = left hippocampal T2 signal; LHV = left hippocampal volume; LTG = lamotrigine; mTL CPS = typical mesial temporal lobe complex partial seizure; PHT = phenytoin; SG TCS = secondary generalized tonic-clonic seizure; TOP = topiramate; VPA = sodium valproate; WAIS-R = revised Weschler Adult Intelligence Scale.

strongly with pre-operative verbal learning (pre-operative immediate list recall), P=0.005 (Fig. 1A). The first principal component derived from the pre-operative memory scores ('pre-operative verbal memory') correlated strongly with pre-operative immediate list recall ($R^2=0.72$, P=0.002) and with pre-operative delayed list recall ($R^2=0.65$, P=0.005), but not with pre-operative story recall ($R^2=0.017$). Similarly, the first principal component derived from the pre-operative—post-operative memory change scores ('post-operative verbal memory change') correlated strongly with pre-operative—post-operative change in immediate list recall ($R^2=0.97$, P<0.001) and with pre-operative—post-operative change in delayed list recall ($R^2=0.38$, P=0.05) but not with pre-operative—post-operative change in story recall ($R^2=0.051$).

Prediction of memory outcome by structural imaging parameters

Post-operative verbal memory change was strongly predicted by left hippocampal volume ($R^2 = 0.53$, P = 0.017) (Fig. 1B).

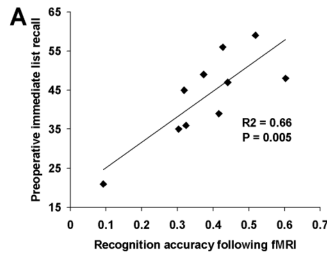
Prediction of memory outcome by pre-operative memory scores

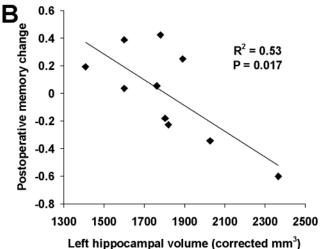
Post-operative verbal memory change was predicted by preoperative verbal memory ($R^2 = 0.50$, P = 0.021) (Fig. 1C).

Prediction of memory outcome by functional imaging parameters (figure 2)

At the group level, a single mesial temporal region showed a correlation between left–right encoding activity difference and post-operative verbal memory change. This region corresponded to the left hippocampus [(-36-20-22), Z=3.64, uncorrected P<0.001, small-volume corrected P=0.016]. Across the whole brain, no other regions survived the threshold chosen.

From a clinical perspective, the power of any predictive test is most relevant at the single subject level. Thus, on an individual subject basis we examined sensitivity, specificity and positive predictive value of pre-operative fMRI in predicting memory decline. We took the simplest approach to this by defining a 'normal' test result as predominant left hippocampal activation [as we previously reported for normal subjects (Richardson et al., 2003)] and an 'abnormal' test result as predominant right hippocampal activation [as we previously reported for patients with more severe left HS (Richardson et al., 2004)]. We defined memory outcome either as 'decline' in memory test score or as 'no decline'. We used two different thresholds to show decline: either any decline greater than zero, or as a decline greater than one standard deviation from the group baseline values. We used a 2×2 contingency-table approach to examine the ability of an 'abnormal' test result to predict 'decline' (Table 2).





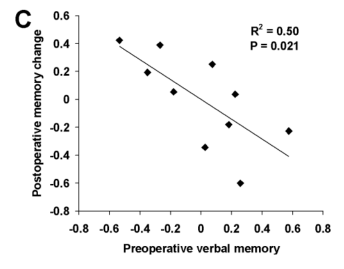


Fig. 1. (A) Correlation between recognition accuracy following fMRI (D') and pre-operative immediate list recall. (B) Correlation between left hippocampal volume (corrected for total intracranial volume) and post-operative memory change (1st principal component of verbal memory change measures; see Methods). (C) Correlation between pre-operative verbal memory (1st principal component of pre-operative verbal memory measures; see Methods) and post-operative memory change (1st principal component of verbal memory change measures; see Methods).

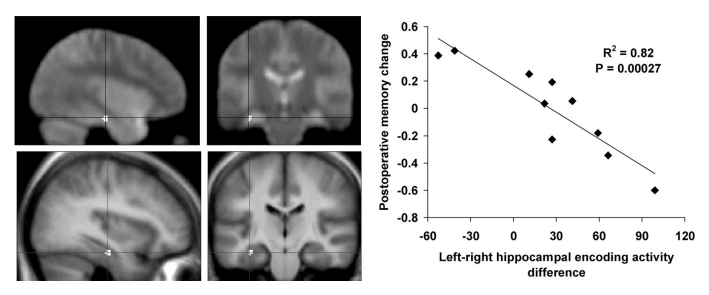


Fig. 2 Voxels showing a significant correlation (P < 0.001) between left-homotopic right voxel value and post-operative memory change (1st principal component of verbal memory change measures; see Methods) mapped onto average T2* image from all subjects (upper left) and average T1 image from all subjects (lower left). The correlation at the peak voxel is illustrated graphically on the right.

Table 2 Sensitivity, specificity and positive predictive value of fMRI for the prediction of pre-operative post—operative change in memory score

	Sensitivity (%)	Specificity (%)	Positive predictive value (%)
Cut-off: decline greater			
than zero from baseline	values		
Immediate list recall	100	100	100
Delayed list recall	100	50	89
Story recall	100	20	56
Cut-off: decline greater			
than I SD from baseline	values		
Immediate list recall	100	40	63
Delayed list recall	100	100	100
Story recall	100	40	63

Prediction of memory outcome: model optimization by stepwise linear regression

Three predictor variables were entered into a model with 'post-operative verbal memory change' as the dependent variable: (i) left-right hippocampal encoding activity difference; (ii) pre-operative verbal memory; and (iii) left hippocampal volume. This model predicted post-operative verbal memory change ($R^2 = 0.92$, P < 0.001). Stepwise linear regression showed only left—right hippocampal encoding activity difference made a significant contribution (significance R^2 change, P > 0.1 for the other variables).

To put this into a clinical context, we constructed an analogous model to predict pre-operative—post-operative change in immediate list recall (using left–right hippocampal encoding activity difference, pre-operative immediate list

recall and left hippocampal volume as predictors). This model predicted pre-operative—post-operative change in immediate list recall ($R^2 = 0.84$, P = 0.008). In this model, stepwise linear regression again showed that only left–right hippocampal encoding activity difference made a significant contribution ($R^2 = 0.74$, P = 0.001).

Finally, we constructed a model to predict pre-operative—post-operative change in delayed list recall (using left-right hippocampal encoding activity difference, pre-operative delayed list recall and left hippocampal volume as predictors). This model predicted pre-operative-post-operative change in delayed list recall ($R^2 = 0.58$, P = 0.13). As in the other models, stepwise linear regression again showed only left-right hippocampal encoding activity difference made a significant contribution ($R^2 = 0.55$, P = 0.015).

Discussion

In this study, we confirmed previous findings that left hippocampal volume and pre-operative verbal memory function predict the extent of verbal memory decline in right-handed subjects with left hippocampal sclerosis undergoing left anterior temporal lobe resection because of intractable MTLE. Our novel finding is that relatively greater verbal memory encoding activity in left hippocampus compared with right hippocampus—as measured using fMRI—predicts the extent of verbal memory decline in the same subjects. Importantly, fMRI was by far the strongest independent predictor of memory decline and a powerful predictor of outcome for individual patients. We show these effects for two different standard clinical measures of verbal memory and even more strongly for a derivative (first principal component) of a range of standard clinical measures, which best accounts for the variability in measures within the patients studied. This is the first time fMRI has shown a predictive value in a clinical setting over and above currently used tests.

Two functional imaging studies have shown prediction of memory outcome in patients undergoing temporal lobe resection for intractable MTLE. One study used blood-flow PET, a technique not generally available in clinical settings and, due to its poor temporal resolution, incapable of resolving activity associated with single events (Henke et al., 2003). Furthermore, only three of their subjects had left MTLE (the group in whom verbal memory decline is anticipated) and none of these experienced any decline in verbal memory. Hence, the study by Henke and colleagues primarily addressed non-verbal memory decline, which is less disabling and less easy to demonstrate, using a method which is not transferable to the clinical environment. This study, like the study we present here, did not address the issue of whether a change in memory test scores is reflected by a subjective change in memory ability which is symptomatic for the patient.

A second study, reported in abstract form, also included a mixed group of eight left and three right MTLE patients (Casasanto et al., 2001). These subjects underwent a task requiring explicit encoding of visual scenes presented in a block design, with a repeated 'scrambled' image as the baseline condition; this design does not allow subsequent memory effects to be revealed. However, the authors undertook a recognition memory test following scanning; a similar visual scene encoding and recognition memory test was also undertaken following subsequent anterior temporal lobe resection. Asymmetry of activation between ipsilateral and contralateral sides in a region of interest (ROI) at the boundary between the hippocampal formation and the lingual gyrus was determined. This asymmetry was strongly correlated with change in recognition memory score between the pre-operative and post-operative assessments. This study therefore also primarily addressed non-verbal memory. In it, memory improvement was seen following surgery in several patients, emphasizing the non-disabling nature of non-verbal memory change in many such patients. In the same group of patients, asymmetry of fMRI activity in the same ROI showed similar efficiency to the IAP for determining seizure outcome (Killgore et al., 1999).

Functional MRI is now a widely used methodology in basic neuroscience, but has yet to show it can provide information useful in clinical settings. Pre-operative localization of motor function is regularly undertaken in some centres to aid neurosurgery in the vicinity of the motor cortex. Although it is often asserted that such an approach provides for a better outcome, there are few supportive data. There is evidence that fMRI of motor function correlates with the sites of motor function determined using electrical stimulation of the cortex during surgery, but the correlation is often imperfect, subject to artefacts in fMRI data, difficult to integrate with other imaging during surgery and possibly requiring subjective interpretation (Stapleton *et al.*, 1997; Krings *et al.*, 2001, 2002; Roux *et al.*, 2001; Liu *et al.*, 2003).

In particular, many brain regions may be activated, especially in patients with lesions, and there is no means to determine which of a number of brain regions activated during an fMRI study is necessary and sufficient for normal function (Krings *et al.*, 2002; Baciu *et al.*, 2003). One study showed a strong prediction from pre-operative data of an immediate post-operative motor deficit following resection of medial frontal lesions if tissue activated in the medial frontal cortex during fMRI was resected, but there was no correlation with outcome a few weeks or months later (Krainik *et al.*, 2001).

Mapping cortex responsible for language function in patients has shown a very strong correlation with IAP findings in the same subjects (Binder et al., 1996; Benson et al., 1999; Springer et al., 1999; Lehericy et al., 2000), although single subject studies are insufficiently sensitive to do more than lateralize language to one hemisphere and more detailed localization has been uncertain. The optimal choice of language task during fMRI remains unclear. An alternative may be to identify the sum total of all regions activated across a range of language tasks (Rutten et al., 2002). In a similar patient group to that presented here, fMRI of a language task showed a predictive value of similar magnitude to the IAP for confrontation naming deficit following left temporal lobe resection (Sabsevitz et al., 2003), but the authors did not present data to show that fMRI was a significantly better predictor of naming outcome compared with IAP.

Our findings have immediate relevance in the evaluation of MTLE patients for possible left anterior temporal lobe resection. There are important further avenues to explore, particularly patients with bilateral pathology and pathology other than hippocampal sclerosis, in whom prediction of verbal memory decline may be difficult. We anticipate that the robust prognostic data provided by our fMRI approach will stimulate the development of further valuable clinical tools utilizing fMRI.

Acknowledgements

We wish to thank Mr William Harkness, neurosurgeon, Dr Maria Thom, histopathologist, and the radiographers of the National Society for Epilepsy MRI Unit for their contributions to this study. We also wish to thank Dr Sanjay Sisodiya and Professor Ley Sander for referring patients. M.P.R. is supported by the Medical Research Council, J.S.D. by the National Society for Epilepsy and R.J.D. by the Wellcome Trust.

References

Baciu M, Le Bas JF, Segebarth C, Benabid AL. Presurgical fMRI evaluation of cerebral reorganization and motor deficit in patients with tumors and vascular malformations. Eur J Radiol 2003; 46: 139–46.

Benson RR, FitzGerald DB, LeSueur LL, Kennedy DN, Kwong KK, Buchbinder BR, et al. Language dominance determined by whole brain functional MRI in patients with brain lesions. Neurology 1999; 52: 798–809.

Binder JR, Swanson SJ, Hammeke TA, Morris GL, Mueller WM, Fischer M, et al. Determination of language dominance using functional MRI: a comparison with the Wada test. Neurology 1996; 46: 978–84.

Cabeza R, Nyberg L. Imaging cognition II: an empirical review of 275 PET and fMRI studies. J Cogn Neurosci 2000; 12: 1–47.

- Casasanto DJ, Glosser G, Killgore WD, Siddiqi F, Falk M, Roc A, et al. Presurgical fMRI predicts memory outcome following anterior temporal lobectomy. J Int Neuropsychol Soc 2001; 8: 183.
- Coughlan A, Hollows S. The Adult Memory and Information Processing Battery. Leeds (UK): Psychology Department, St James' Hospital; 1985.
- Crawford PM. Epidemiology of intractable focal epilepsy. In: Oxbury JM, Polkey CE, Duchowny M, editors. Intractable focal epilepsy. London: W.B. Saunders; 2000. p. 25–40.
- Detre JA, Maccotta L, King D, Alsop DC, Glosser G, D'Esposito M, et al. Functional MRI lateralization of memory in temporal lobe epilepsy. Neurology 1998; 50: 926–932.
- Duncan JS, Bartlett P, Barker GJ. Technique for measuring hippocampal T2 relaxation time. AJNR Am J Neuroradiol 1996; 17: 1805–10.
- Dupont S, Van de Moortele PF, Samson S, Hasboun D, Poline JB, Adam C, et al. Episodic memory in left temporal lobe epilepsy: a functional MRI study. Brain 2000; 123: 1722–32.
- Friston KJ, Holmes AP, Worsley KJ, Poline JB, Frith CD, Frackowiak RSJ. Statistical parametric maps in functional imaging: a general linear approach. Hum Brain Mapp 1995; 2: 189–210.
- Friston KJ, Fletcher P, Josephs O, Holmes A, Rugg MD, Turner R. Eventrelated fMRI: characterizing differential responses. Neuroimage 1998; 7: 30–40
- Golby AJ, Poldrack RA, Illes J, Chen D, Desmond JE, Gabrieli JD. Memory lateralization in medial temporal lobe epilepsy assessed by functional MRI. Epilepsia 2002; 43: 855–63.
- Hauser AW. Incidence and prevalence. In: Engel H Jr, Pedley TA, editors. Epilepsy: a comprehensive textbook, Vol. 1. Philadelphia: Lippincott-Raven; 1998. p. 47–57.
- Helmstaedter C, Elger CE. Cognitive consequences of two-thirds anterior temporal lobectomy on verbal memory in 144 patients: a three-month follow-up study. Epilepsia 1996; 37: 171–80.
- Henke K, Treyer V, Weber B, Nitsch RM, Hock C, Wieser HG, et al. Functional neuroimaging predicts individual memory outcome after amygdalohippocampectomy. Neuroreport 2003; 14: 1197–202.
- Hermann BP, Wyler AR, Somes G, Berry AD, Dohan FC. Pathological status of the mesial temporal lobe predicts memory outcome from left anterior temporal lobectomy. Neurosurgery 1992; 31: 652–6.
- Jokeit H, Ebner A, Holthausen H, Markowitsch HJ, Moch A, Pannek H, et al. Individual prediction of change in delayed recall of prose passages after left-sided anterior temporal lobectomy. Neurology 1997; 49: 481–7.
- Jokeit H, Okujava M, Woermann FG. Memory fMRI lateralizes temporal lobe epilepsy. Neurology 2001; 57: 1786–93.
- Killgore WD, Glosser G, Casasanto DJ, French JA, Alsop DC, Detre JA. Functional MRI and the Wada test provide complementary information for predicting post-operative seizure control. Seizure 1999; 8: 450–5.
- Krainik A, Lehericy S, Duffau H, Vlaicu M, Poupon F, Capelle L, et al. Role of the supplementary motor area in motor deficit following medial frontal lobe surgery. Neurology 2001; 57: 871–8.
- Krings T, Reinges MH, Erberich S, Kemeny S, Rohde V, Spetzger U, et al. Functional MRI for presurgical planning: problems, artefacts, and solution strategies. J Neurol Neurosurg Psychiatry 2001; 70: 749–60.
- Krings T, Topper R, Willmes K, Reinges MH, Gilsbach JM, Thron A. Activation in primary and secondary motor areas in patients with CNS neoplasms and weakness. Neurology 2002; 58: 381–90.
- Lee GP, Park YD, Westerveld M, Hempel A, Blackburn LB, Loring DW. Wada memory performance predicts seizure outcome after epilepsy surgery in children. Epilepsia 2003; 44: 936–43.

- Lehericy S, Cohen L, Bazin B, Samson S, Giacomini E, Rougetet R, et al. Functional MR evaluation of temporal and frontal language dominance compared with the Wada test. Neurology 2000; 54: 1625–33.
- Lepage M, Habib R, Tulving E. Hippocampal PET activations of memory encoding and retrieval: the HIPER model. Hippocampus 1998; 8: 313–22.
- Liu H, Hall WA, Truwit CL. The roles of functional MRI in MR-guided neurosurgery in a combined 1.5 Tesla MR-operating room. Acta Neurochir Suppl 2003; 85: 127–35.
- Loring DW, Meador KJ, Lee GP, King DW, Nichols ME, Park YD, et al. Wada memory asymmetries predict verbal memory decline after anterior temporal lobectomy. Neurology 1995; 45: 1329–33.
- Richardson MP, Strange BA, Duncan JS, Dolan RJ. Preserved verbal memory function in left medial temporal pathology involves reorganization of function to right medial temporal lobe. Neuroimage 2003; 20 Suppl 1: S112–9.
- Richardson MP, Strange BA, Dolan RJ. Encoding of emotional memories depends on amygdala and hippocampus and their interactions. Nat Neurosci 2004; 7: 278–85.
- Roux FE, Ibarrola D, Tremoulet M, Lazorthes Y, Henry P, Sol JC, et al. Methodological and technical issues for integrating functional magnetic resonance imaging data in a neuronavigational system. Neurosurgery 2001; 49: 1145–56; discussion 1156–7.
- Rutten GJ, Ramsey NF, van Rijen PC, Noordmans HJ, van Veelen CW. Development of a functional magnetic resonance imaging protocol for intraoperative localization of critical temporoparietal language areas. Ann Neurol 2002; 51: 350–60.
- Sabsevitz DS, Swanson SJ, Hammeke TA, Spanaki MV, Possing ET, Morris GL, 3rd, et al. Use of pre-operative functional neuroimaging to predict language deficits from epilepsy surgery. Neurology 2003; 60: 1788–92.
- Sass KJ, Westerveld M, Buchanan CP, Spencer SS, Kim JH, Spencer DD. Degree of hippocampal neuron loss determines severity of verbal memory decrease after left anteromesiotemporal lobectomy. Epilepsia 1994; 35: 1179–86.
- Springer JA, Binder JR, Hammeke TA, Swanson SJ, Frost JA, Bellgowan PS, et al. Language dominance in neurologically normal and epilepsy subjects: a functional MRI study. Brain 1999; 122: 2033–46.
- Squire LR. Memory and the hippocampus: a synthesis from findings in rats, monkeys and humans. Psychol Rev 1992; 99: 195–231.
- Squire LR, Zola-Morgan S. The medial temporal lobe memory system. Science 1991; 253: 1380–6.
- Stapleton SR, Kiriakopoulos E, Mikulis D, Drake JM, Hoffman HJ, Humphreys R, et al. Combined utility of functional MRI, cortical mapping, and frameless stereotaxy in the resection of lesions in eloquent areas of brain in children. Pediatr Neurosurg 1997; 26: 68–82.
- Strange BA, Henson RN, Friston KJ, Dolan RJ. Brain mechanisms for detecting perceptual, semantic, and emotional deviance. Neuroimage 2000; 12: 425–33.
- Trenerry MR, Jack CR, Ivnik RJ, Sharbrough FW, Cascino GD, Hirschorn KA, et al. MRI hippocampal volumes and memory function before and after temporal lobectomy. Neurology 1993; 43: 1800–5.
- Tulving E. Memory and consciousness. Can Psychol 1985; 26: 1-12.
- Van Paesschen W, Sisodiya S, Connelly A, Duncan JS, Free SF, Raymond AA, et al. Quantitative hippocampal MRI and intractable temporal lobe epilepsy. Neurology 1995; 45: 2233–40.
- Wiebe S, Blume WT, Girvin JP, Eliasziw M. A randomized, controlled trial of surgery for temporal-lobe epilepsy. N Engl J Med 2001; 345: 311–8.