Transient global amnesia: implicit/explicit memory dissociation and PET assessment of brain perfusion and oxygen metabolism in the acute stage

Francis Eustache, Béatrice Desgranges, Marie-Christine Petit-Taboué, Vincent de la Sayette, Véronique Piot, Christelle Sablé, Gilles Marchal, Jean-Claude Baron

Abstract

Objective—To assess explicit memory and two components of implicit memory—that is, perceptual-verbal skill learning and lexical-semantic priming effects—as well as resting cerebral blood flow (CBF) and oxygen metabolism (CMRO₂) during the acute phase of transient global amnesia.

Methods—In a 59 year old woman, whose amnestic episode fulfilled all current criteria for transient global amnesia, a neuropsychological protocol was administered, including word learning, story recall, categorical fluency, mirror reading, and word stem completion tasks. PET was performed using the [¹⁵O] steady state inhalation method, while the patient still exhibited severe anterograde amnesia and was interleaved with the cognitive tests.

Results—There was a clear cut dissociation between impaired long term episodic memory and preserved implicit memory for its two components. Categorical fluency was significantly altered, suggesting word retrieval strategy—rather than semantic memory—impairment. The PET study disclosed a reduced CMRO₂, with relatively or fully preserved CBF in the left prefrontotemporal cortex and lenticular nucleus, and the reverse pattern over the left occipital cortex.

Conclusions—The PET alterations with patchy CBF-CMRO₂ uncoupling would be compatible with a migraine-like phenomenon and indicate that the isolated assessment of perfusion in transient global amnesia may be misleading. The pattern of metabolic depression, with sparing of the hippocampal area, is one among the distinct patterns of brain dysfunction that underlie the (apparently) uniform clinical presentation of transient global amnesia. The finding of a left prefrontal hypometabolism in the face of impaired episodic memory and altered verbal fluency would fit present day concepts from PET activation studies about the role of this area in episodic and semantic memory encoding/retrieval. Likewise, the changes affecting the lenticular nucleus but sparing the caudate would be consistent with the normal performance in perceptual-verbal skill learning. Finally, unaltered lexical-semantic priming effects, despite left temporal cortex hypometabolism, suggest that these processes are subserved by a more distributed neocortical network.

Keywords: transient global amnesia; implicit/explicit memory dissociation; oxygen metabolism; PET

Forty years after its original description, the pathophysiology of transient global amnesia remains elusive, with three major hypotheses: a transient ischaemia, a seizure process, or a migraine phenomenon. Likewise, despite extensive neurological accounts, only 35 cases in whom formal (although usually rather limited) neuropsychological assessment was performed during the acute stage have been reported. Although these studies concur in showing that transient global amnesia is a pure syndrome of isolated memory impairment akin to the permanent amnesic syndrome (PAS), with a consistent dissociation between preserved short term but impaired long term memory (LTM), the exact profile of memory impairment in transient global amnesia remains incompletely known.

Thus only a few studies have assessed the distinct components of LTM during episodes of transient global amnesia, notably implicit memory (skill learning and priming effects). Apart from the above mentioned impairment in episodic memory, which is intrinsic to the definition of transient global amnesia, one consistent finding has been that semantic memory is preserved. Although a dissociation between declarative memory and skill learning is suggested by the well known finding that patients with transient global amnesia are occasionally able to perform complex previously acquired skills, preserved acquisition of new skills has been formally documented only rarely. Likewise for priming effects, only perceptual priming effects have been studied whereas lexical-semantic priming effects have remained unexplored thus far.

Studies of resting cerebral perfusion and metabolism during transient global amnesia are of special interest because they may shed light not only on the pathophysiology, but also on the mechanisms of the cognitive impair-
ment of transient global amnesia. Unfortunately, only few such studies have been possible for obvious logistical reasons. Only three patients have had both perfusion and metabolism assessed with PET, two reported in abstract form and one as a full report. In the rest solely brain perfusion was assessed, by means of SPECT. Although many of these studies (performed either in the acute stage, the early recovery phase, or immediately after the episode) not unexpectedly disclosed unilateral or bilateral alterations in perfusion or metabolism in mesial temporal, or thalamic regions, several did find occasionally isolated changes in neocortical, notably frontal and temporal, regions, either unilaterally or bilaterally. These unexpected imaging findings made the important suggestion that an acute impairment of episodic memory could be subtended by different localisations of synaptic dysfunction. Memory was concurrently investigated only in a few of the above studies as part of either a restricted, or a more extensive assessment, but in all the imaging modality used was SPECT, and almost none provided a detailed interpretation in terms of the mechanisms of transient amnesia. As uncoupling between CBF and metabolism is frequent in acute brain conditions, perfusion may not constitute a valid index of synaptic function under these circumstances, and thus SPECT may provide incomplete or even misleading information for both the pathophysiology and the neurobiological basis of memory dysfunction during transient global amnesia.

Our aims were to formally investigate both memory (notably perceptual-verbal skill learning and lexical-semantic priming effects), and resting cerebral perfusion and oxygen metabolism with PET during the acute phase of transient global amnesia, according to a specially designed prospective protocol. We anticipated that, by carrying out both types of investigations concurrently, we would gain further insight into the cognitive and physiological pattern(s) of transient global amnesia.

**Subjects and methods**

**PATIENTS**

Over a period of six months, four patients with suspected transient global amnesia were seen by us in the emergency unit of the University Hospital of Caen. Three fulfilled the operational criteria for transient global amnesia set out by Caplan and modified by Hodges and Warlow, but in two of them, the episode was over before the tests could be completed. In the last patient, a 59 year old housewife, it was possible to perform both the neuropsychological investigation and PET during the episode; this patient forms the basis of the present article.

**CONTROLS**

A group of seven subjects, strictly matched to the patient in sex, age (mean (SD): 59.4 (2.8) years), and educational level was used for normative neuropsychological data, whenever these were unavailable from the literature, or for personal versions or French adaptations of published tests. Selection was made based on lack of neurological or psychiatric disorder or general cognitive impairment (as assessed with Signoret’s “Batterie d’Efficience cognitive” (BEC), in which each score has to be above 9/12 (see below)). Likewise, for the PET study, we constructed a control group made of 10 optimally healthy, unmedicated subjects of mean age matched to that of the patient (mean (SD): 56.5 (5.8) years). In both groups, the minimum educational level was equivalent to “certificat d’études primaires”—a now abandoned diploma obtained after eight years of primary education.

**NEUROPSYCHOLOGICAL INVESTIGATION**

**Protocol**

We prospectively constructed a brief modular neuropsychological protocol such that it could be administered in sequences interleaved with both the emergency medical investigations required to evaluate acute amnesia (for example, CT, ECG, EEG), and the PET study.

The major aim of this protocol was to assess separately the two components of implicit memory—that is, priming effects and skill learning (see below). In addition, explicit episodic memory and other cognitive functions were assessed with subtests of the “Batterie d’Efficience Mnésique” (BEM) and of the BEC respectively. For general cognitive functions we assessed mental control, visuoconstructive abilities (with the copy of three geometrical figures, two simple and a complex one), orientation, problem solving, and semantic similarities. Semantic memory was more specifically assessed by means of both categorical verbal fluency (generating as many names of animals as possible in one minute) and naming of 12 verbalisable (high frequency) line drawings. For episodic memory, we assessed (1) visual episodic memory with the immediate free recall of the Signoret’s complex geometric figure and the free recall and the recognition of six verbalisable line drawings (different from the set of 12 used in the naming test); and (2) verbal episodic memory, by means of story recall and word learning tasks. In the word learning task, the order of recall (which was left without specific instruction) made it possible to calculate the primacy and recency effects, using a method modified from Tulving and Colotla who consider a word to be recalled from short term memory (STM) if its presentation is separated from its recall by at most six words, and from LTM otherwise. Our modified procedure uses the subject’s own verbal span instead of the fixed number six. In addition to the recency effect, STM was also assessed with the forward verbal span.

For implicit memory, we elected to restrict our time constrained protocol to the study of perceptual-verbal procedural memory and lexical-semantic priming effects, using modified versions of the mirror reading and the word stem completion paradigms respectively.

The mirror reading paradigm was adapted from Cohen and Squire. The mirror words were presented in 20 pairs in a 20 page
The priming effect could give rise to at least 10 common words. That each stem (both primed and distractors) length (six to eight letters), and in such a way 17000 occurrences per 100 million) and match the primed words and the distractors for allowed us to examine the effect of learning in each session by comparing the mean slope of each regression line obtained across the seven control subjects with zero. Likewise, retention of the mirror reading skill from session I to session II across subjects was assessed by comparing the formula (B-A)/(B+A) with zero, where A and B are the y intercepts of the regression lines for sessions I and II, respectively.

Lexical-semantic priming effects were assessed with a modified word-stem completion paradigm. In the first part (study phase), and without having been instructed to memorise, the subject was required to make a sentence from each word of a series of 15 written words (of which the 10 middle words were later used for assessment of priming effects; the first two and last three being present only to reduce primacy and recency effects). During the second part (test phase), given immediately after the first, 20 written three letter stems were presented to the subject, who had to complete them orally with the first word that came to mind. Ten of the stems were the beginning of non-primed words that matched the selected words in the notebook. The number of primed words minus the number of non-primed words that matched the selected distractors.

**Statistical analysis**

The patient’s performance in each subtest of Signoret’s batteries (the BEC and the BEM) was assessed against their published normative data, when available; for each subtest, a z score was calculated and judged significant if beyond the one tailed t value for P<0.05 corresponding to the sample of control subjects for each test—that is, 1.70 for the BEC (n=30 subjects) and 1.72 for the BEM (n=21 subjects). For the scores obtained in the subtests of the BEM and the BEC for which no normative data are available (namely, forward verbal span, primacy, and recency effects) or the subtests for which we used personal adaptations (copy of complex geometric figure), as well as for the word stem completion and mirror reading tasks, we used as normative data the performances of our group of seven age matched and education matched controls; statistical assessment was as above, except that the t value was 1.94 (P<0.05, n=7). For the mirror reading task, the patient’s slopes and retention index were compared with the control subjects according to the same general statistical procedure.

**PET investigation**

**PET imaging**

The PET study was done according to a standardised procedure. We used the “O steady state inhalation method” with serial arterial “O counting and transmission attenuation correction. We used a high resolution, 7 slice tomograph (TTV03, LETI, Grenoble, France), with final apparent image resolution ~11 × 11 × 9 mm, xyz. We used the stereotaxic procedure derived from Fox et al., based on the glabella and the inion bony landmarks. The seven PET planes were made parallel to the glabella-inion line and lay from −4 mm to +68 mm relative to it. To restrain the head during scanning, we used a Laitinen stereotaxic frame. Furthermore, head position during scanning was continuously monitored and any displacement immediately corrected, by laser beams projected on forehead ink marks. The raw “O and “O images were systematically realigned with the dedicated software of Woods et al. We obtained parametric images of cerebral blood flow (CBF; ml/100 g/min), oxygen extraction fraction (OEF, dimensionless), and cerebral metabolic rate of oxygen (CMRO2; ml/100/ min), corrected for intravascular radiotracer, by applying to the raw PET images, pixel by pixel, a set of non-linear equations.

**Regions of interest (ROI) procedure**

A three dimensional MRI (made of 128 contiguous T1 weighted scans) was performed one week after the PET (GE Advantage 1.5 T), according to the SPGR procedure. By means of dedicated software enabling MRI data acquired in three dimensions to be resliced to the PET coordinates with respect to the glabella-inion line landmarks (which has known relations with the anterior commissure—posterior commissure reference points), circular regions of interest (ROIs; n=103, radius=7mm) were positioned directly over the MRI cross sections corresponding to the PET planes,
according to Talairach’s stereotaxic atlas, to sample neocortical, limbic (especially hippocampal), subcortical (namely, caudate nucleus, lentiform nucleus, and thalamus,) and cerebellar areas. These ROIs were projected on to corresponding PET matrices, and the ROI pixel values were averaged according to anatomical-functional cortical and non-cortical areas to yield regional values for CBF, OEF, and CMRO2. (Right-left percentage differences for homologous regions (asymmetry indices) were also calculated.)

Statistical assessment
We compared the regional index values and the (R-L) asymmetry indices obtained for each brain area in our patient to single subject confidence limits (mean (SD)\textsubscript{t,p,df}) calculated from data obtained with the same methodology in 10 healthy volunteers of similar mean age. Because of the different number of ROIs included in a given anatomical-functional area (for example, from 2 for the thalamus to 14 for the lateral-frontal cortex), the accuracy in regional index estimation varied and the confidence intervals differed widely from one region to another. In addition, the confidence limits for the (R-L) asymmetry indices could be weighted towards the right or the left side due to underlying overall asymmetries in controls.

Results
Figure 1 shows the time course of the different events related to the transient global amnesia episode.

CASE HISTORY
The patient is a 100% right-handed 59-year-old woman with a history of treated arterial hypertension, but otherwise healthy. There was no histology of migraine.

On 16 June 1994, at 7:45 am, her husband was awakened by noises of retching from his wife, who also complained of headache. She asked her husband repeatedly about the place and date. There was no accompanying motor, sensory, or visual impairment, clouding of consciousness, speech deficit, behavioural impairment, or convulsion. At 9:00 am, her general practitioner visited her at home. He noted memory impairment and temporal disorientation; blood pressure was 150/90 mmHg. It was noted that the patient did not remember her sister’s move to another nearby city 13 years before. During the remainder of the day, repetitive queries were noted by both her husband and her sister as well as by the hospital staff. Although no precipitating event was evident, the patient had felt tired for several days before the episode, and two potentially emotional circumstances existed: the prospect of being awarded a poetry prize the following day, and the unexpected death of her favourite poet and singer the day before.

At 12 noon, the patient was seen at the emergency ward of the university hospital of Caen. Language, behaviour, and somatic neurological examination were unremarkable. Unenhanced emergency CT of the head was normal, as were also the standard EEG and the ECG. At 4:00 pm, a neurologist noted an anterograde amnesia with temporal disorientation as well as a retrograde amnesia: the patient could not recall any of her personal events of the day, or of the day before (shopping in Bayeux, a walk at Omaha Beach). The patient had also forgotten about both the European Assembly elections (on 12 June 1994) and the just finished and locally vastly publicised 50th celebration of D Day (6 June 1994). She was then sent for PET, which started at 6:30 pm and was completed by 8:00 pm (fig 1). Neuropsychological examination was done just

Figure 1. Time course of the different events related to the transient global amnesia episode. The neuropsychological assessment was performed in three parts: part I: explicit memory, word stem completion, general battery; part II: mirror reading (first session); part III: mirror reading (second session).
Table 1 Performances of patient and matched control subjects (mean (SD)) in cognitive non-nemonic functions and explicit memory tests

<table>
<thead>
<tr>
<th>Test</th>
<th>Patient Controls</th>
<th>Raw score</th>
<th>z Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>General cognitive functions:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mental control‡</td>
<td>12 (0)</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Copy of Signoret’s complex geometrical figure†</td>
<td>11.9 (0.24)</td>
<td>12</td>
<td>+4.42</td>
</tr>
<tr>
<td>Copy of geometrical figure‡</td>
<td>10.73 (1.14)</td>
<td>10</td>
<td>+4.64</td>
</tr>
<tr>
<td>Orientation‡</td>
<td>11.5 (0.86)</td>
<td>12</td>
<td>+4.58</td>
</tr>
<tr>
<td>Problem solving and semantic similarities‡</td>
<td>11.27 (0.87)</td>
<td>12</td>
<td>+0.84</td>
</tr>
<tr>
<td>Semantic memory:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Categorical verbal fluency‡</td>
<td>11.53 (0.86)</td>
<td>8</td>
<td>−4.10</td>
</tr>
<tr>
<td>Naming‡</td>
<td>10.9 (0.96)</td>
<td>12</td>
<td>+1.14</td>
</tr>
<tr>
<td>Visual episodic memory:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Six verbalisable line drawings (free recall and recognition)</td>
<td>10.93 (1.01)</td>
<td>8</td>
<td>−2.90</td>
</tr>
<tr>
<td>Recall of complex geometrical figure†</td>
<td>8.57 (1.24)</td>
<td>3</td>
<td>−4.49</td>
</tr>
<tr>
<td>Verbal episodic memory:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Story recall‡</td>
<td>8.48 (1.09)</td>
<td>4.5</td>
<td>−3.65</td>
</tr>
<tr>
<td>Word learning global score†</td>
<td>8.31 (1.25)</td>
<td>4.5</td>
<td>−3.05</td>
</tr>
<tr>
<td>Primacy effect‡</td>
<td>20.57 (5.09)</td>
<td>6</td>
<td>−2.86</td>
</tr>
<tr>
<td>Short term memory:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Word learning recency effect‡</td>
<td>3.71 (2.87)</td>
<td>6</td>
<td>+4.80</td>
</tr>
<tr>
<td>Forward verbal span†</td>
<td>5.1 (1.07)</td>
<td>7</td>
<td>+1.77</td>
</tr>
<tr>
<td>Controls</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* P < 0.01; ** P < 0.001 reduced performance v controls (see methods for calculations).
† From Signoret’s “Batterie d’efficience cognitive”.
‡ From Signoret’s “Batterie d’efficience mnésique”.
§ Normative data from control sample of seven subjects (see methods for details).
The maximal possible score for all tests was 12, except for the primacy and recency effects, and for the forward verbal span.

before and after PET scanning, while the patient was fully alert, and confirmed the existence of a major impairment of episodic memory (see next section). Thus the PET study was completed while the patient still exhibited severe anterograde amnesia.

At 1100 pm, after completion of the neuropsychological protocol, the patient still exhibited both anterograde and retrograde amnesia, although she clearly had entered the recovery phase. For instance, she still could not recall the death of her favourite poet-singer the day before. By the next morning, she was clinically normal. Four formal follow up neuropsychological examinations were performed, one and 14 days and four and 13 months after the episode (see below).

A three dimensional MRI, performed on 12 July, was normal, including both a detailed assessment of the hippocampus (with specially oriented cuts) and T2 weighted imaging. A cervical Doppler ultrasound investigation was also negative. A neurological examination at six months follow up was unremarkable. As of August 1996, her health has been unremarkable. She has a memory blank (amnestic gap) of the entire day.

NEUROPSYCHOLOGICAL EXAMINATION DURING THE ACUTE STATE

General assessment (table 1)

Mental control, copy of geometric figures, orientation, problem solving, semantic similarities, naming, and forward verbal span were not significantly altered (all scores were above the respective control means). Conversely, verbal fluency was significantly impaired (reduced output of correct items without errors nor perseverations), as were the verbal and the visual components of episodic memory. Regarding the word learning task, in addition to a significantly low global score, she exhibited an impaired primacy effect but a preserved recency effect. Furthermore, the order of recall, which reflects strategy, was totally inconsistent across trials (data not shown). However, perseverations or overt frontal signs were not found.

Skill learning task (fig 2 and table 2)

Figure 2 and table 2 show the learning curves and corresponding quantitative values (slope 1, slope 2, and retention index) respectively. Figure 2 shows that the control subjects learned the procedure rapidly during the first session, and showed no further learning at the second session. Our patient’s data were not different and showed a better performance after learning relative to the control mean. In controls, the mean slope (on log transformed data) for the first session was significantly different from zero, indicating a significant learning effect (table 2). The corresponding slope for the patient was not significantly different from that of controls. For the second session, the slope in controls was not different from zero and showed a pronounced intersubject variability. The corresponding slope in the patient fell well within the normal range. The retention index was significantly different from zero in controls (indicating significant retention of the procedure from the first to the second session), and showed no alteration in the patient.

Word stem completion (table 3)

The patient’s performances in this task did not fall outside the 95% normal confidence limits for any score, and especially for priming effects.

FOLLOW UP NEUROPSYCHOLOGICAL ASSESSMENT

Follow up neuropsychological assessments purposefully focused on the functions which were altered during the episode, employing more formal tests whenever relevant. The morning after the episode, an IQ of 110 was obtained on Raven’s progressive matrices46 and an MQ of 107 on the Wechsler memory scale (with normal scores in all the subtests). Word fluency, assessed with the same task as in the acute stage protocol, had normalised (13 and 22 names of animals in one and two minutes respectively). The performance in the copy of Rey’s complex figure was normal (34/36), but its delayed recall after three minutes was still poor (7/36, z = −2.5, P<0.05).

Two weeks after the episode, the global mnemonic score assessed with the BEM144 was strictly normal at 100.5, and verbal fluency was again normal. Four months after the episode, the patient was still clinically normal, the MQ was 101 and all subtests scores were normal, as
was verbal fluency; however, performance in the delayed recall of Rey’s figure was unchanged (6/36). With the BEC, there was normal performance in both free recall (6/6) and recognition (6/6) of verbalisable line drawings, tests which were both significantly impaired in the acute stage (table 1). Thirteen months after the transient global amnesia episode, the neuropsychological examination was normal, including recall and recognition of the simple, non-verbalisable line drawings of the BEM, except for the delayed recall of Rey’s and Signoret’s complex geometric figures. Her poor performance in the recall of complex figures would reflect inadequate visuospatial abilities before the transient global amnesia episode, which the patient herself rated so since childhood. Furthermore, this peculiarity concerned preferentially the long term memory for complex geometric figures, but spared the delayed free recall of both verbalisable and simple non-verbalisable line drawings.\textsuperscript{49,50}

**PET STUDY**

Appreciable head movements did not occur during the entire PET session; neither were movement artefacts evident on PET OEF images. Figure 3 shows the set of PET images with coregistered MRI. Visual interpretation suggested a reduction in both the CMRO\textsubscript{2} and OEF over the left, compared with the right cortical convexity. The CMRO\textsubscript{2} appeared especially reduced in the left frontal and temporal regions, as well as over the left lenticular nucleus. For CBF, there appeared a moderate reduction in the left, compared with the right lateral and posterior occipital cortex. The hippocampal area appeared unremarkable.

These findings from visual interpretation was confirmed by the objective ROI analysis; table 4 shows the main results. Although globally in the low range, the CMRO\textsubscript{2} was significantly reduced relative to age matched controls only in the left lateral frontal cortex; no regional value for CBF or OEF fell outside the confidence limits from control data. The CMRO\textsubscript{2} for the whole left neocortex was reduced relative to the right, the L/R\% asymmetry failing just short of significance (right = 1.95 ml/100 g/min; left=1.79 ml/100 g/min; L/R=0.915, P=0.06). Regionally, the CMRO\textsubscript{2} was significantly reduced in the left compared with the right lateral frontal cortex, lateral temporal cortex, and lentiform nucleus, with a similar although non-significant trend for the thalamus. An analysis of the individual cortical gyri to assess in further detail the distribution of metabolic abnormalities disclosed significant CMRO\textsubscript{2} reductions in the left, compared with the right, middle, and inferior lateral frontal gyri and inferior lateral temporal cortex (data not shown). Relative to the right side, the OEF was reduced in the left lateral frontal cortex and lentiform nucleus (P<0.05 for both), as well as in the left lateral temporal and lateral parietal cortex (P=0.06

![Figure 2](http://jnnp.bmj.com/)

**Table 3** Word stem completion: results for control subjects (mean (SD)) and patient

<table>
<thead>
<tr>
<th>Controls (n=10*)</th>
<th>Patient</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primed words</strong></td>
<td></td>
</tr>
<tr>
<td>4.7 (1.8)</td>
<td>2</td>
</tr>
<tr>
<td><strong>Non-primed words</strong></td>
<td>0</td>
</tr>
<tr>
<td>3.7 (1.7)</td>
<td>2</td>
</tr>
<tr>
<td><strong>Priming effect</strong></td>
<td></td>
</tr>
<tr>
<td>1 (0.8)</td>
<td>-1.2 (NS)</td>
</tr>
<tr>
<td>1 (0.7)</td>
<td>-1 (NS)</td>
</tr>
</tbody>
</table>

* Normative data from Desgranges et al.\textsuperscript{46,47} Presentation is otherwise similar to table 1. Each maximum possible score is 10.

\textsuperscript{49} The hippocampal area appeared unremarkable.

\textsuperscript{50} These findings from visual interpretation was confirmed by the objective ROI analysis; table 4 shows the main results. Although globally in the low range, the CMRO\textsubscript{2} was significantly reduced relative to age matched controls only in the left lateral frontal cortex; no regional value for CBF or OEF fell outside the confidence limits from control data. The CMRO\textsubscript{2} for the whole left neocortex was reduced relative to the right, the L/R\% asymmetry failing just short of significance (right = 1.95 ml/100 g/min; left=1.79 ml/100 g/min; L/R=0.915, P=0.06). Regionally, the CMRO\textsubscript{2} was significantly reduced in the left compared with the right lateral frontal cortex, lateral temporal cortex, and lentiform nucleus, with a similar although non-significant trend for the thalamus. An analysis of the individual cortical gyri to assess in further detail the distribution of metabolic abnormalities disclosed significant CMRO\textsubscript{2} reductions in the left, compared with the right, middle, and inferior lateral frontal gyri and inferior lateral temporal cortex (data not shown). Relative to the right side, the OEF was reduced in the left lateral frontal cortex and lentiform nucleus (P<0.05 for both), as well as in the left lateral temporal and lateral parietal cortex (P=0.06

**Table 4** PET results

<table>
<thead>
<tr>
<th>CBF (ml/100 g min)</th>
<th>CMRO\textsubscript{2} (ml/100 g min)</th>
<th>OEF</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Right</strong></td>
<td><strong>Left</strong></td>
<td><strong>L/R %</strong></td>
</tr>
<tr>
<td>Lateral frontal cortex</td>
<td>29</td>
<td>28</td>
</tr>
<tr>
<td>Lateral temporal cortex</td>
<td>29</td>
<td>28</td>
</tr>
<tr>
<td>Lateral parietal cortex</td>
<td>24</td>
<td>26</td>
</tr>
<tr>
<td>Occipital cortex\textsuperscript{‡}</td>
<td>31</td>
<td>26</td>
</tr>
<tr>
<td>Hippocampus</td>
<td>29</td>
<td>28</td>
</tr>
<tr>
<td>Thalamus</td>
<td>37</td>
<td>30</td>
</tr>
<tr>
<td>Lentiform nucleus</td>
<td>36</td>
<td>29</td>
</tr>
</tbody>
</table>

\* p < 0.05, \*\* p < 0.01 with respect to confidence intervals calculated in 10 age matched normal controls (see methods).

† P = 0.06; \‡ this region includes the lateral occipital cortex and the cuneus. CBF = cerebral blood flow; CMRO\textsubscript{2} = cerebral metabolic rate of oxygen; OEF = oxygen extraction fraction.
for both), whereas the CBF was significantly reduced in the left lateral occipital cortex. For the hippocampus, there was not even a trend for changes in any of the variables assessed (table 4).

Over and above these findings, which essentially confirmed the visual interpretation, the ROI analysis disclosed the following additional significant changes (all \( P < 0.05 \)): (1) a significant increase in the OEF in the left occipital cortex, (2) a significant increase in CBF in the left parietal cortex, and (3) a significant decrease in CBF in the left lentiform nucleus. Not shown in table 4, there was a lack of significant changes in the caudate nucleus and the cerebellum.

**Discussion**

This is the first study to report on a patient with transient global amnesia fulfilling the operational criteria for transient global amnesia of Hodges and Warlow,\(^2\) examined in the acute stage with a prospective protocol comprising both a brief neuropsychological assessment (including tests of skill learning and priming effects) and PET. Nausea and headache occur at times during transient global amnesia.\(^2\)\(^5\) No epileptic feature was noted at any time and there was no history of head injury. All medical investigations, including EEG, CT, and detailed MRI, were strictly normal, and no further episode has occurred with a follow up over two years.

**NEUROPSYCHOLOGICAL DATA**

The patient’s neuropsychological assessment confirmed the presence of severe anterograde amnesia and showed a significant impairment of episodic LTM (for both verbal and visual components), whereas STM was strictly intact. The preserved recency effect but altered primacy effect, showing a clear cut dissociation between LTM and STM, is consistent with most previous transient global amnesia case reports (see introduction). The patient’s categorical fluency was also significantly reduced.
during the episode but had normalised as early as the morning after. Impaired verbal fluency (either categorical or formal) has been previously reported during transient global amnesia, although this certainly is not a consistent finding.\textsuperscript{5,10-11} Categorical fluency deficits can result from a perturbation of either semantic memory or retrieval strategy. The preservation of all other language functions in our patient (namely, spontaneous verbal expression and comprehension, confrontation naming, and semantic similarities) favours the integrity of semantic stores as also supported by the normality of lexical-semantic priming effects. This interpretation would also fit well with Hodges\textsuperscript{11} finding of intact semantic memory in his two patients. Thus although we cannot formally exclude the existence of a mild semantic impairment, the reduction of verbal fluency in our patient most likely reflected an alteration in strategy, which would account for the observed inconsistency in recall order across trials in the word learning test.

In our study, we used two distinct paradigms to assess skill learning and priming effects. Both these components of implicit memory were preserved. Using identification of fragmented digits, Goldenberg \textit{et al}\textsuperscript{21} were the first to claim preservation of perceptual skill learning in transient global amnesia. Kazui \textit{et al}\textsuperscript{18} reported preservation of cognitive, perceptual-motor, and perceptual-verbal procedural memory, using the Tower of Toronto, a drawing skill, and reading skill tests respectively; however, the last did not involve the acquisition of a new skill and therefore would not be rated by most as a procedural memory test. Thus the present study seems to be the first to have formally assessed perceptual-verbal skill learning during transient global amnesia. We used the mirror-reading paradigm, which is well recognised to assess this process. Both the acquisition and the retention of this skill were similar to control subjects (see fig 2; note that the better performance of control subjects in the first trial of session II may reflect an involvement of declarative memory,\textsuperscript{12} which was impaired in the patient).

For priming effects, we used an original version of the word stem completion paradigm, specifically designed to investigate lexical-semantic priming effects, which did not fall outside the (admittedly wide\textsuperscript{3}) normal range. Recently, both Kazui \textit{et al}\textsuperscript{18} and Kapur \textit{et al}\textsuperscript{24} showed preservation of perceptual priming in three and one cases, respectively. Overall, therefore, we document for the first time the preservation of both perceptual-verbal skill learning and lexical-semantic priming effects in transient global amnesia.

\textbf{PET FINDINGS}

A PET study was carried out while the patient was still severely amnesic, as documented by the cognitive tests performed during and after scanning. All the changes were confined to the left cerebral hemisphere, although with different patterns of CBF-CMRO\textsubscript{2} uncoupling from region to region. Specifically, there was a metabolic depression maximum in the left frontal cortex (especially affecting the middle and inferior frontal gyri), left temporal cortex (especially inferior), and left lentiform nucleus. In these regions, the CBF was fully preserved in the former two and reduced, (but less so than the CMRO\textsubscript{2}) in the third region respectively, with consequently a reduction in the OEF in all three regions. Other abnormalities included (1) a mildly increased CBF with reduced OEF but normal CMRO\textsubscript{2} in the left parietal cortex, and (2) a reduction in CBF with increased OEF, again with normal CMRO\textsubscript{2}, in the left occipital cortex. There was a non-significant trend for reduced CBF and CMRO\textsubscript{2} in the left thalamus. Finally, the hippocampal area exhibited not even a trend for changes in any variable; this does not reflect a lack of sensitivity of our methodology to changes in the hippocampal region, as shown previously by, for example, age related reductions in hippocampal CMRO\textsubscript{2}, correlated with verbal episodic memory declines.\textsuperscript{30,34}

This report of predominantly neocortical changes with unaffected hippocampal physiology during transient global amnesia, although at first sight surprising, is not isolated. As stated in the introduction, several previous functional neuroimaging studies including our previous PET report,\textsuperscript{23} have reported neocortical hypoperfusion/hypometabolism, in some instances associated with ipsilateral or bilateral thalamic changes, yet without hippocampal changes. Although the last may have occurred at earlier times into the episode, the patient was severely amnesic during PET scanning. Thus the present report further strengthens the idea that, at least in some patients, the clinical expression of a global but transient amnesia may correspond to predominantly neocortical dysfunction; this distribution of physiological changes in transient global amnesia is in variance with the topography of brain damage in the PAS, despite a very similar neuropsychological pattern. This discrepancy may in part reflect the single major difference between transient global amnesia and PAS—namely, that the latter usually represents the final, stable phase of an initially more diffuse and complex neuropsychological picture; in other words, the clinical expression of a given acute focal brain dysfunction may considerably overstate that which permanently endures once brain reorganisation has taken place.

Compared with the right side, the relative preservation (or even increase) of CBF with mildly but significantly decreased OEF in the left frontal, temporal, and parietal cortices would suggest a flow metabolism uncoupling compatible with, for example, prior ischaemia, prior seizure, or prior spreading depression.\textsuperscript{35-37} Previously, Volpe \textit{et al}\textsuperscript{20} also reported a reduced OEF in the hippocampal region of their patient with transient global amnesia; however, in the case of Baron \textit{et al},\textsuperscript{21} the OEF was normal. Thus different pathophysiological mechanisms may trigger transient global amnesia, or, alternatively, each patient is “snapshot” with PET at different phases of one (and the same) process. The presence of mild but significant...
Transient global amnesia: implicit/explicit memory dissociation

Changes with reduced CBF and increased OEF in the left compared with the right occipital cortex is of great interest as it suggests the mechanism underlying transient global amnesia in this case also included oligemia as one of its expressions. The aura phase of classic migraine is characterised by a narrow wave of hyperperfusion slowly spreading from the occipital cortex towards the forebrain and leaving in its aftermath a prolonged and extensive hypoperfusion with (in the single instance of protracted aura captured by PET) locally increased OEF. These changes have been largely interpreted as reflecting the human equivalent of the phenomenon of spreading depression of Leão. The nature and complex pattern of associated abnormalities in CBF, OEF, and CMRO₂ in our patient would be compatible with these migrainous phenomena, and spreading depression has been speculated previously as one trigger of transient global amnesia. From a clinical point of view, there are many arguments which point towards a link between transient global amnesia and migraine, both epidemiological and symptomwise and our present findings would add a pathophysiological argument as one further support to this hypothesis.

The present findings also clearly illustrate that in the acute phase of transient global amnesia the CBF may be uncoupled from CMRO₂ (in either direction). Thus perfusion imaging studies without concomitant assessment of metabolism (as with SPECT) may mislead the interpretation in terms of synaptic function.

NEUROPSYCHOLOGY–METABOLISM RELATIONS

Impaired episodic memory

Although both clinical and animal neuropsychology have shown the implication of the hippocampal complex in episodic memory, the CMRO₂ in the hippocampal region was not significantly altered in our patient, despite concurrent massive episodic amnesia. As stated above, this finding is, however, consistent with several reports, although hippocampal changes were effectively detected in others. Furthermore, as PET or SPECT in other conditions (for example, normal aging, PAS, Alzheimer’s disease) have documented significant relations between hippocampal area metabolism and episodic memory deficits, the imaging technology is not to be called into question. PET activation studies in normal subjects did not systematically show involvement of the hippocampus in episodic memory tasks. The implication of the hippocampus in episodic memory is probably much more complex than previously thought and may not be systematically required. It is worth recalling in this context that a typical PAS may develop despite an intact hippocampus if the lesion affects other parts of the episodic memory (limbic) network—for example, the diencephalic area. In the present patient, there was a trend for left thalamic hypometabolism which however did not reach significance. In the case reported by Baron et al., right sided tandem prefrontal thalamic hypometabolism was found.

It is tempting, even if speculative, to relate the deficit in episodic memory in our patient with hypometabolism in the left prefrontal cortex. Thus although subjects with frontal lobe impairment do not exhibit genuine PAS, they do have difficulties in making temporal order judgements (frequency and recency assessment), increased susceptibility to interference, impaired free recall, but preserved recognition and “source” amnesia. Furthermore, PET activation studies have recently clarified the role of the prefrontal cortex in memory. Thus according to the hemispheric encoding retrieval asymmetry (HERA) model, the left and right prefrontal cortices are involved in the operations of encoding and retrieval of episodic information respectively. It is of note that the hypometabolic zone in the left prefrontal cortex in our patient affected the inferior frontal gyrus, precisely where Shallice et al. found a significant activation during episodic memory encoding in normal subjects. Our PET results would thus be consistent with an encoding deficit; however, we did not specifically evaluate encoding versus retrieval of new episodic information. In the single detailed PET report on transient global amnesia published to date, a right prefrontal hypometabolism was found; unfortunately, a neuropsychological assessment could not be concurrently performed. SPECT reports have occasionally mentioned either right or left prefrontal hypoperfusion during transient global amnesia, but encoding versus retrieval was not assessed. Thus the hypothesis of a causal relation between prefrontal hypometabolism and episodic memory impairment during transient global amnesia will have to be tested in future work. However, it is unclear whether the left prefrontal hypometabolism in our patient also accounts for the presence of retrograde amnesia, which must reflect a retrieval impairment; however, this retrieval mechanism may well differ from the one operative in episodic tasks assessing anterograde amnesia, as suggested by dissociations in cases of pure retrograde amnesia.

IMPAIRED VERBAL FLUENCY

During the episode, our patient exhibited an impaired categorical verbal fluency. This was the only significant cognitive impairment apart from the deficit of episodic memory. It is likely that this impairment was related to the left prefrontal or temporal hypometabolism disclosed by PET. Previously, Goldberg had reported a patient with impaired verbal fluency together with left frontal hypoperfusion during a transient global amnesia episode. Aphasic patients with left temporal damage exhibit anomia, and several PET activation studies have localised semantic representations for words and objects in the left temporal lobe. However, as discussed above, the neuropsychological data were more consistent with an impairment in verbal fluency from strategy rather than semantic knowledge alteration, which would point towards the left frontal lobe. Accordingly, left prefrontal cortex activations...
have been consistently found across various PET paradigms involving either categorical verbal fluency or a word generation paradigm. It is thought that the left cortex is specifically engaged in the search, manipulation, and mental organisation of semantic information, either automatic or willed, including retrieval from semantic stores. This function of the left dorsolateral prefrontal cortex is incorporated in the HERA model (elaborated by Tulving et al. and supported by Shallice et al.).

INTACT PERCEPTUAL-VERBAL SKILL LEARNING

The pattern of metabolic depression observed in the patient, sparing the cerebellum and the caudate, would be compatible with preserved perceptual-verbal skill learning. Thus neuropsychological studies suggest a contribution of the cerebellum and of a corticostriatal system in skill learning. Furthermore, caudate nucleus lesions impair the acquisition of mirror reading but not target pursuit skills, and putaminal lesions the reverse pattern. PET activation studies in healthy subjects have attributed a major role to the cortical-striatal-cerebellar motor system in motor skill learning. To our knowledge, however, no activation study has so far specifically focused on perceptual-verbal skill learning.

PRESERVED LEXICAL-SEMANTIC PRIMING EFFECTS

Because a selective impairment in priming effects as a result of brain damage seems at best very rare, the neural substrates of this intriguing phenomenon remain largely unknown to date. However, the fact that priming effects are intact in PAS indicates that the limbic system (hippocampus, diencephalon, anterior cingulate cortex) is not required for this function. Lexical-semantic priming is impaired in Alzheimer’s disease suggesting that widespread lesions are required in association with other cognitive deficits. Our patient exhibited left temporal cortex hypometabolism that was matched by a preserved pattern of metabolic depression observed in the patient, sparing the cerebellum and the caudate, would be compatible with preserved perceptual-verbal skill learning. Thus neuropsychological studies suggest a contribution of the cerebellum and of a corticostriatal system in skill learning. Furthermore, caudate nucleus lesions impair the acquisition of mirror reading but not target pursuit skills, and putaminal lesions the reverse pattern. PET activation studies in healthy subjects have attributed a major role to the cortical-striatal-cerebellar motor system in motor skill learning. To our knowledge, however, no activation study has so far specifically focused on perceptual-verbal skill learning.

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