Low-dose Propofol-induced Amnesia Is Not due to a Failure of Encoding

Left Inferior Prefrontal Cortex Is Still Active

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Background: Propofol may produce amnesia by affecting encoding. The hypothesis that propofol weakens encoding was tested by measuring regional cerebral blood flow during verbal encoding.

Methods: Seventeen volunteer participants (12 men; aged 30.4 ± 6.5 yr) had regional cerebral blood flow measured using H2015 positron emission tomography during complex and simple encoding tasks (deep vs. shallow level of processing) to identify a region of interest in the left inferior prefrontal cortex (LIPFC). The effect of either propofol (n = 6, 0.9 µg/ml target concentration), placebo with a divided attention task (n = 5), or thiopental at sedative doses (n = 6, 3 µg/ml) on regional cerebral blood flow activation in the LIPFC was tested. The divided attention task was expected to decrease activation in the LIPFC.

Results: Propofol did not impair encoding performance or reaction times, but impaired recognition memory of deeply encoded words 4 h later (median recognition of 35% [interquartile range, 17–54%] of words presented during propofol vs. 65% [38–91%] before drug; P < 0.05). Statistical parametric mapping analysis identified a region of interest of 6.6 cm2 in the LIPFC (T = 7.44, P = 0.014). Regional cerebral blood flow response to deep encoding was present in this region of interest in each group of volunteers; however, during propofol infusion, only the propofol group continued to have borderline significant activation in this region (T = 4.00, P = 0.063).

Conclusions: If the amnesic effect of propofol were solely due to effects on encoding, activation in the LIPFC should be minimal. Because LIPFC activation was not totally eliminated by propofol, the amnesic action of propofol must be present in other brain regions and/or affect other memory processes.

ONE of the still unsolved mysteries of anesthesia is how temporary amnesia is produced during wakefulness by drugs such as midazolam or propofol. Dense amnesia, even for intense events such as intraoperative awakening, has been demonstrated for propofol. The memories most sensitive to propofol are those that are consciously accessible in rich detail and include contextual information such as when and where they occurred.2,3 These memories are encoded and retrieved by the episodic memory system, which includes the hippocampus as the central, key component of memory processing.4 Episodic memory can be conceptualized as the processes of encoding, storing, and then retrieving information. Although testing recognition memory some time after the administration of propofol clearly demonstrates the presence of amnesia, one cannot determine which memory process or processes are affected.2 Imaging brain activity using electroencephalography, positron emission tomography (PET), or functional magnetic resonance imaging during behavioral paradigms that isolate specific memory processes can characterize the brain processes underlying specific components of episodic memory function.5–7 These neuroimaging techniques may reveal differences in the processes and neural structures underlying a given behavior (e.g., encoding) in the setting of matched behavioral performance, as has been done in the elderly versus the young.8,9 Specifically, at similar performance levels, the elderly recruit additional brain regions during encoding, and do not demonstrate event-related potentials in the left prefrontal cortex as large as do the young during semantic retrieval and encoding operations. That is, behavioral measures alone are insufficient to fully characterize memory processes.10,11 The current study focuses on the processes and neural structures that underlie encoding, using PET imaging of cerebral blood flow (CBF) to more fully characterize the effects of propofol, if any, on this key memory process.

Encoding is defined as the process of acquiring and representing information in memory. For the purposes of the current study, memory processes after encoding are divided into working and long-term memories. Encoding is closely related to working memory, and overlapping brain regions are involved.12–15 Working memory is defined as a short-term retention of information no longer available in the environment, and is of limited capacity (e.g., 7 ± 2 items).12,16 After approximately 18 s, information can only be retained in working memory by ongoing rehearsal.17 Long-term memory is defined here as memory that is not working memory, i.e., memory retained longer than 20 s after encoding in the absence of rehearsal. There are a number of phases of

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long-term memory, which we do not differentiate for the purposes of the current study. 18

One method that allows assessment of the effects of a drug on working and long-term memory processes as defined above is the continuous recognition task. We previously showed that at low doses, propofol did not impair encoding or working memory as measured by behavioral responses on the continuous recognition task, but still produced amnesia at the end of the study day.19 Despite normal encoding performance by behavioral measures in the presence of propofol, the processes and neural structures underlying this behavior might still be affected by propofol. We have recently presented suggestive evidence that event-related potential measures of recognition memory in the presence of propofol are altered even when performance is unchanged.20 Similar changes by propofol of brain processes underlying encoding could potentially affect longevity of memories over time. Such subtle effects on verbal memory, for example, have been demonstrated using transcranial magnetic stimulation.21 22

One brain region important in verbal encoding is the left inferior prefrontal cortex (LIPFC). A behavioral paradigm that isolates encoding activity and reveals this brain region as important in this aspect of episodic memory is the levels of processing manipulation (LoP). This task involves evaluations of a series of words using either semantic, i.e., meaning-based, or stimulus-based processing.23–25 In this task, words could be categorized as representing a living object, or whether the word was spoken in a male or female voice, for example. Semantic evaluations invoke more complex memory operations that contact already stored knowledge. These operations result in deeply encoded memories that are enhanced and better retained than those formed during shallow, stimulus-based processing. Memory processes underlying encoding are isolated by this task as evaluation occurs on the first presentation of the stimulus. These are revealed by the comparison of brain images obtained during deep versus shallow categorizations. The LIPFC and, in some studies, the hippocampus are identified using this methodology.8 24 26 Because the LoP improvement in memory is preserved in the presence of benzodiazepines, albeit at a lower level, this manipulation can be used to index encoding processes in the presence of doses of drug that produce memory impairment.27 28

As the dose of propofol increases, associated sedation occurs.2 The difference between an amnesic dose of propofol without and one with associated sedation is small and not predictable in advance in a given individual. Behavioral measures reveal that higher, sedative doses of propofol can interfere with working memory and thereby impair encoding.19 Therefore, in the current study, it is important to assess the effects of attention and sedation on the brain processes underlying encoding to clearly interpret the effects of propofol on these.

A behavioral manipulation that influences encoding via attentional mechanisms is the divided attention task. This task requires the participant to pay attention to a diversionary task when an encoding task is performed. The need for divided attention during encoding decreases subsequent recall or recognition memory.20 Importantly for the current study, impaired encoding from divided attention is associated with decreased brain activation.30–34 Therefore, this task was included in the placebo group in the current study to isolate attentional effects in the LIPFC during encoding. Likewise, to assess the influence of sedation on LIPFC responses to the levels of processing manipulation, thiopental was included as an additional control group because it interferes with memory principally by sedation.2

The hypothesis tested in this study is that amnesic doses of propofol weaken encoding, as measured by effects on the regional cerebral blood flow (rCBF) response to the levels of processing manipulation in the LIPFC region of the prefrontal cortex. Our study is designed to closely parallel that of Kapur et al.,24 who were able to isolate LIPFC activity using this paradigm. A dose of propofol was chosen to produce amnesia but not sedation.2 19 35 Behavioral measures of sedation were obtained by measuring accuracy of word categorizations and associated reaction times during the encoding tasks. Amnesic effects were assessed by recognition of words at various times after encoding, including at the end of the study day. The divided attention and thiopental groups served as controls for attentional and sedation effects on imaged brain responses. As explained below, fewer numbers of participants completed the study than anticipated. Therefore, the primary response measure to address the hypothesis is the effect size of brain activation in the LIPFC, as estimated by T values of deep versus shallow encoding brain image contrasts in the study groups.

Materials and Methods

Participants

Twenty-five healthy volunteers (19 men) were recruited through newspaper advertisements and were paid for their participation. Participants were right-handed, proficient English speakers with normal hearing, aged between 18 and 45 yr (mean age, 31.0 ± 10.5 yr). Exclusion criteria included use of psychoactive medication; history of recreational drug abuse; head trauma resulting in loss of consciousness; psychiatric, neurologic, cardiovascular, or respiratory disease; major psychiatric disease in a first-degree relative; claustrophobia; carpal tunnel syndrome; allergy to eggs; or acute intermittent porphyria. This investigation was approved by the Hospital Institutional Review Board and Radiation Safety Committees of Memorial Sloan-Kettering Cancer Center and Weill Cornell Medical Center, New York,

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New York. Informed consent was obtained in writing before accrual of participants took place.

 Procedures

 Levels of Processing Tasks. The LoP paradigm was used to encode auditory stimuli into verbal memory. In the deep encoding task, participants performed a semantic categorization task. Immediately after hearing each word, participants categorized it as representing either a living or a nonliving object. Choices were communicated to the investigators with a button press response with the right hand. Likewise, during shallow encoding, participants categorized the voice speaking the word as male or female.

 Before Study Day. During an orientation session, detailed information was given on study procedures, including radiation dose. Tests of handedness (Edinburgh Handedness Inventory) and vocabulary (vocabulary subtest of Wechsler Adult Intelligence Scale–Revised) were administered, followed by a brief physical examination. Participants were screened for normal hearing using the Coren-Hakstian Hearing Screening Inventory before recruitment into the study. Inform consent was obtained at that time.

 Study Day Procedures.

 Preparation. On the study day, participants arrived at approximately 8 AM, being nil per os since midnight. A 64-channel electroencephalographic cap (Compumedics, Neuroscan, El Paso, TX) was applied using standard procedures. Electroencephalographic data are not reported here. Venous catheters were inserted before transfer to the PET suite. Dextrose, 5%, ½ normal saline, at approximately 100 ml/h was administered intravenously. All participants practiced the divided attention task and were informed that they might need to perform it after the start of drug infusion. However, they were not informed that the task was only used in the placebo group.

 Behavioral Tasks. Participants served as their own controls in the three study groups, those being propofol, placebo with a divided attention task (placebo/DA), or thio.

 PET Imaging in Relation to Behavioral Tasks. The participant’s total time in the PET scanner was approximately 2.5–3 h, with scans obtained every 12 min (table 1). The interval was longer just before drug infusion was started, to allow for a brief break for the participant and to allow establishment of the target drug concentration after the start of drug infusion.

 The baseline condition consisted of scanning during shallow and deep encoding tasks. After this, two recognition scans were needed to image the first recognition task of baseline words. The reason for needing two scans over one recognition task was as follows. One scan was obtained during presentation of old words presented during the previous deep encoding task, whereas the other was obtained during presentation of new words. The order of old and new word scans was counterbalanced across participants. Each recognition scan was immediately preceded and followed by presentation of words of the opposite category than those presented during imaging, so that to the participants, each recognition scan involved the same ratio of old and new words. Therefore, two recognition scans comprise one recognition task for the full set of deeply encoded words in both baseline and drug conditions.

 Drug infusion was started after the first recognition task, and then a second recognition task again comprising two PET scans of the baseline encoded words was obtained, to assess the effect of the presence of drug on recognition of words encoded in the absence of drug. After this second recognition task, shallow and deep encoding tasks/scans in the drug condition were obtained, followed by two more sets of recognition scans, to match the baseline condition. Drug infusion was then stopped.

 End-of-day Recognition Task. Approximately 2.5 h after PET scanning and drug infusion were finished, a series of four recognition tasks (end-of-day recognition [EODr]g) was given in randomized order, in which participants heard words previously presented (old words) during both shallow and deep encoding tasks in baseline and drug conditions, along with an equal number of distractor words (new words) not heard before. Participants made a yes/no decision regarding whether they had heard the word during PET scanning. Recognition...
tion tasks were administered at approximately 286 min after deep encoding in the baseline condition and approximately 207 min after deep encoding in the drug condition. Participants were discharged home after meeting standard criteria for discharge for ambulatory surgery.

Because of scheduling issues, eight volunteers underwent structural magnetic resonance imaging scanning late in the afternoon in the same facility after lunch but before delayed recognition testing, whereas nine returned for a magnetic resonance imaging scan on a separate occasion.

Materials

O-15 Administration and PET Brain Images. For each PET scan, approximately 10 mCi H$_2^{15}$O was delivered intravenously at a constant rate over 20 s via an infusion pump. Scans were obtained on a GE Advance scanner (GE Medical Systems, Waukesha, WI) in the three-dimensional “septa out” mode. The resolution of the PET camera in this mode is approximately 5.2 mm in all dimensions. Imaging started with the arrival of radioactivity at the brain. Three 30-s frames were obtained during each scan, corresponding to the highest rate of uptake of tracer into the brain. Two computed tomography scans were obtained before baseline and drug conditions to correct for attenuation of signal in its passage through bone and cerebral tissue and to align the PET camera to each volunteer’s brain. The images were reconstructed using filtered back projection and standard clinical protocols and were stored as “counts” images (counts of coincidence events, expressed as nCi/ml). The structural (T1) magnetic resonance scan for each participant was used in coregistration and spatial normalization in statistical analysis of brain images.

Study Drugs. Placebo solutions had the same appearance as thiopental (multivitamin solution in saline) or propofol (intraalipid). Both investigators and participants were blinded as to the nature of the study drug.

Computer-controlled Drug Infusion. Study drug was given by intravenous infusion using STANPUMP software controlling a Harvard22 infusion pump (Harvard Apparatus, Holliston, MA). This software produces a constant plasma and/or effect site concentration based on population pharmacokinetics adjusted for weight, age, and sex depending on the particular kinetic data set used. Propofol was infused using the Schnider kinetic software. The data set. Target concentrations were chosen based on previous experience so that propofol would have some degree of memory impairment, with thiopental being given at an equivalent sedative dose. The target concentrations used were 0.9 µg/ml propofol or 3 µg/ml thiopental.$^{2,19,35}$

Auditory Stimuli. Auditory stimuli were presented using Stim software (Compumedics, Neuroscan, Charlotte, NC). A total of 60 words were presented over a 3-min time period. Word presentation started 30 s before PET imaging began and continued for 60 s after scanning was completed. Behavioral results reflect the performance on the total set of words presented. Participants wore foam earphones inserted into the auditory canal (EarLink Auditory Systems, Indianapolis, IN). Words were delivered every 3 s at 80 dB sound pressure level. After insertion of earphones and positioning in the scanner, participants were tested for clear and bilaterally equal perception of words before imaging was started.
Word Stimuli. Words presented in a given deep encoding task were presented two more times during PET scanning before final EODrcg. Shallowly encoded words were only presented once (during the encoding task) before final EODrcg (table 1). Separate word lists were prepared for baseline and drug conditions and were presented in counterbalanced order across participants. Two-syllable words from the Toronto Word Pool (word frequency < 100; mean duration 766 ms) were used. Care was taken to avoid using similar-sounding words as targets and distractors. Words were digitized for computer presentation as female or male voices. Words for the deep encoding task (living/nonliving) were presented in a third, neutral voice.‡‡ At EODrcg, the stimuli were presented in the same voice used at encoding.

Divided Attention Stimuli. Participants randomly assigned to the placebo/DA condition heard a series of tones randomly interspersed with word stimuli in the drug condition only. Tones and words were presented by two separate computers so that intervals between words and tones were truly random. Tones were of either 1,000- or 1,100-Hz frequency, 800-ms duration, presented randomly with the high-pitched tones occurring at an overall frequency of 30%. Upon hearing the high-pitched tone, participants pushed a button with the left hand, while still using the right hand for responses to encoding and recognition tasks. All participants in all study groups held both response devices.

Statistical Analysis

In planning of the current study, power analysis of rCBF response determined that nine participants in each study group would be required to achieve 90% power at \( \alpha = 0.01 \) based on an effect size estimated at 1.5. Unanticipated technical issues with O-15 production resulted in smaller accruals than this desired goal. Therefore, the power to accept or reject the hypothesis as stated is limited, and results are to be interpreted in light of this consideration.

Behavioral Data. Performance accuracy and reaction times on the encoding tasks during PET imaging were obtained as measures of sedation in the drug as compared with the baseline conditions. EODrcg was the primary response variable in testing the memory effects of study manipulations. However, interim memory performance was also assessed using recognition tasks of deeply encoded words with the participant in the PET scanner at approximately 18 and 42 min after deep encoding.


Final recognition tasks were administered at the end of the study day. All recognition scores were corrected for false alarms before statistical analysis. False alarms are defined as a new word (distractor) that the participant identified as old (having been presented during PET imaging). Because of the small number of participants, nonparametric Kruskal-Wallis tests for independent samples for the main effect of group (propofol, placebo/DA, and thiopental) and Friedman tests for related samples were used to test the effect of stimulus type (shallow or deep encoding words) factors. Within-participant changes (by group and condition) were addressed using nonparametric Wilcoxon signed rank tests. Data were combined across groups in the baseline condition in certain comparisons because all participants underwent the same treatment in this condition. A threshold significance of \( P < 0.05 \) was chosen, although \( P \) values near this cutoff are reported as trends because of small sample sizes. Statistical analyses of behavioral data were conducted using SPSS version 12.0 (Chicago, IL).

Statistical Analysis of PET Images. Images were reconstructed into .mnc format for analysis. Statistical analysis was performed using Statistical Parametric Mapping implemented in Pro Matlab version 7.0, release 14 (Mathworks, New York, NY). PET images were coregistered to the participant’s structural T1 weighted magnetic resonance image.

Images were realigned to the first baseline or drug scan and normalized into Montreal Neurologic Institute brain image space with resultant voxel size of \( 3 \times 3 \times 3 \) mm. A 10-mm gaussian smoothing kernel was used to accommodate interpersonal variations in gyral anatomy and facilitate intersubject averaging with a resultant smoothness of \( 12.4 \times 13.2 \times 15.8 \) (x, y, z) mm. Mean global CBF was normalized to 50 ml \( \cdot \) 100\(^{-1} \) g brain tissue \( \cdot \) min\(^{-1} \). Statistical analysis used a proportional scaling model, which allows a differing relation between rCBF effect depending on global CBF.

The main statistical parametric mapping contrast was constructed to identify regions of the brain more active in the deep than in the shallow encoding tasks using images from 17 participants in the baseline condition using a voxel-wise \( P \) value threshold of \( P < 0.001 \) (fig. 1). Normally, a region of interest (ROI) is defined on the basis of \textit{a priori} considerations. In the current study, the paradigm used is almost exactly the same as that used by Kapur et al.\(^{24} \) Rather than using defined anatomical structures, the ROI was critically defined by the contrast of deep \textit{versus} shallow encoding, where all aspects of the task (e.g., auditory stimulation, button press) were identical except for cognitive processing. The ROI thus identified in the current study corresponded closely to the results of Kapur et al. Supporting the use of this ROI as an \textit{a priori} region for statistical analysis is replication of this region in other studies of encoding memory using functional magnetic resonance imaging methods.\(^{13,25,31} \)
Fig. 1. Regions of the brain that were more active in the deep encoding task (semantic, meaning-based evaluation of words: “Did the word represent a living object?”) than in shallow encoding (stimulus-based evaluation of words: “Was the word spoken in a male or female voice?”). Statistical significance in this map represents greater regional cerebral blood flow as quantified using Statistical Parametric Mapping 99 at a voxel-wise P value threshold of \( P = 0.001 \) (\( T = 3.55 \)). The regions displayed were identified from the baseline condition using all 17 subjects in the current study. The region of interest in the left inferior prefrontal cortex (peak activation at Montreal Neurologic Institute \( x, y, z \) coordinates of \( -48, 12, 3 \) mm, \( T = 7.44 \)) has been associated with verbal encoding tasks in multiple previous studies. This region was considered an \textit{a priori} region of interest because the location of activation is similar to that identified by other investigators. A smaller region of activation, not considered as an \textit{a priori} region of interest, was evident near the midline in the anterior cingulate gyrus (peak activation at \( -9, 39, 42 \), \( T = 6.03 \)). The colored bar represents the \( T \) value significance in this comparison of deep greater than shallow encoding.

This ROI thus identified was used in Statistical Parametric Mapping 99 analyses of images in each treatment group in baseline and drug conditions. A small volume correction was applied to correct for multiple comparisons in the ROI, and the measures of significance, \textit{i.e.}, \( T \)-values, of activation of deep \textit{versus} shallow encoding contrasts were used as an estimate of effect size. The deep \textit{versus} shallow encoding contrast in the baseline condition revealed another region of activation present near the midline, but this was not considered an \textit{a priori} ROI.

Neuroanatomical labeling of the ROIs was performed using automated anatomical labeling software, which interfaces with Statistical Parametric Mapping 99.

### Results

#### Participants

Of the 25 recruited, 17 participants (12 men) fully completed the study, with 6 receiving propofol, 5 receiving placebo/DA, and 6 receiving thiopental (table 2). As noted in the Materials and Methods, the relatively small numbers in each group did not achieve the planned accrual. Of the 8 participants excluded, 3 fell asleep (responded to less than 50 of 60 word stimuli) during one of the encoding tasks (2 thiopental, 1 propofol), 1 did not understand instructions, in 1 the scanner malfunctioned, and 3 were assigned to a placebo/no-DA group (not analyzed because of insufficient numbers). Thirteen of the participants were college graduates or had an advanced degree; the remainder had at least some postsecondary education. Mean scores on the Edinburgh Handedness Inventory were strongly right-handed: 88.2 (14.9). Percentile scores on the vocabulary subtest of the Wechsler Adult Intelligence Scale–Revised ranged from 50 to 99, with a mean (SD) of 83.9 (15.9). Eleven participants (64.7%) were white.

#### Behavioral Results

The goal of the experimental task, \textit{i.e.}, the levels of processing paradigm, was to ensure encoding of verbal information into long-term memory in both baseline and drug conditions. Memories had to be of sufficient strength so that recognition would be above chance at the end of the study day. To achieve these results, participants had to perform at high levels in the confines of the PET scanner, which might have been difficult especially in the drug condition. Measures of encoding performance demonstrated that participants performed accurately in both baseline and drug conditions. As anticipated, the semantic categorization task proved to be more difficult than speaker voice categorization as measured by accuracy rates and reaction times. This relation was present in both baseline and drug conditions and resulted in robust memories for deeply encoded words.

#### Encoding Accuracy and Reaction Times

Performance during encoding tasks in both baseline and drug conditions was more than 88% accurate, indicating that attention was given to the task in both study conditions (table 3). There was no significant effect of study group (propofol, placebo/DA, or thiopental) on accuracy in the baseline or drug condition for either stimulus type (Kruskal-Wallis, not significant). Therefore, all groups performed equally well in both conditions. In the baseline condition (\( n = 17 \)), there was a strong effect present for stimulus type (deep \textit{vs.} shallow words) on accuracy of categorization, with shallow categorizations (96 ± 7.6% correct) being more accurate than deep categorizations (90.5 ± 5.1%) (Friedman, \( P = 0.002 \)). In the group by condition comparisons, significant effects were present in propofol baseline and possibly drug conditions (Wilcoxon signed rank, \( P = 0.05 \) and \( P = 0.06 \)), thiopental baseline condition (\( P = 0.05 \)).
and possibly placebo/DA drug condition \((P = 0.08)\), as indicated in table 3.

Also indicative that semantic categorization was a more difficult task were the longer reaction times required to categorize these stimuli as compared with speaker voice determinations. As with encoding accuracy, there was no effect of group (propofol, thiopental, placebo/DA) on reaction times in either drug or baseline conditions for either deep or shallow stimulus types (Kruskal-Wallis, not significant). Therefore, all groups reacted equally fast to categorize stimuli in both conditions. In the baseline condition, there was a strong effect of stimulus type on reaction time, with deep categorizations taking longer (1,410 ± 216 ms) than shallow categorizations (1,097 ± 262 ms) (Friedman, \(P = 0.008\)). In group by condition testing, significantly longer reaction times to deep encoding stimuli were present for placebo/DA baseline condition (Wilcoxon signed rank, \(P = 0.04\)) and thiopental baseline (\(P = 0.05\); table 4).

Paired contrasts of condition effect (baseline, drug) for both performance parameters in each group revealed no significant effects, except possibly for encoding accuracy in the placebo/DA group in the deep encoding task (Wilcoxon signed rank, \(P = 0.06\); all others, not significant). Therefore, in a given group, participants had similar behavioral responses to the categorization tasks regardless of whether drug was present.

### Memory

As revealed in the baseline condition, semantic categorization in the level of processing paradigm was a more difficult task. This resulted in better memory for these stimuli at the end of the study day, as is typical for this paradigm. All recognition scores were corrected for false alarms (new words that were categorized as old) before statistical testing (the false-alarm rates are reported in table 5). In all groups, significantly greater memory at the end of the study day for deeply encoded words was present when compared with shallow ones, in both baseline and drug conditions (Wilcoxon signed rank, \(P < 0.05\) for all EODrcg cells in table 5). Essentially, when corrected for false alarms, recognition scores for shallowly encoded words at EODrcg were zero. Therefore, EODrcg memory for deeply encoded words was above chance levels for all groups in both baseline and drug conditions.

As would be expected by repeated presentations of stimuli in the recognition tasks conducted in the PET scanner soon after encoding, some improvement in memory occurred for deeply encoded words, as represented by differences in recognition 1 (Rcg1) and recognition 2 (Rcg2) correct responses in figure 2. A significant learning effect was present on the second recognition task in the placebo/DA group in both baseline and drug conditions (Wilcoxon signed rank, Rcg1 vs. Rcg2, baseline \(P = 0.043\), drug \(P = 0.043\)) and likely also in the propofol but not thiopental groups in the baseline condition (Wilcoxon signed rank, Rcg1 vs. Rcg2, propofol \(P = 0.058\)).

The key behavioral result in the current study was a significant effect of group (propofol, placebo/DA, thiopental) on recognition of deeply encoded words at the end of the study day (Kruskal-Wallis, \(P = 0.031\)). Using planned comparisons, deeply encoded words in the drug condition for propofol were more poorly recognized than baseline encoded words at the end of the day (Wilcoxon signed rank, \(P = 0.046\)). This differential effect of propofol on memory for deeply encoded words was evident in the second recognition testing approximately 42 min after deep encoding in the drug condition.

### Table 3. Percent Correct Encoding Performance While in the PET Scanner

<table>
<thead>
<tr>
<th>Group</th>
<th>Baseline Shallow Encoding</th>
<th>Baseline Deep Encoding</th>
<th>Drug Shallow Encoding</th>
<th>Drug Deep Encoding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Propofol, n = 6</td>
<td>100 (5.4)</td>
<td>90.0 (9.1)*</td>
<td>97.5 (7.5)</td>
<td>89.2 (11.7)†</td>
</tr>
<tr>
<td>Placebo/DA, n = 5</td>
<td>95.0 (18.3)</td>
<td>91.7 (6.7)</td>
<td>96.7 (9.2)</td>
<td>88.3 (8.4)†</td>
</tr>
<tr>
<td>Thiopental, n = 6</td>
<td>98.3 (1.7)</td>
<td>89.2 (11.2)*</td>
<td>95.9 (17.0)</td>
<td>91.7 (8.8)</td>
</tr>
</tbody>
</table>

Data are shown as median (interquartile range). Shallow encoding accuracy < deep encoding accuracy by Wilcoxon signed rank test.

\* \(P < 0.05\). † \(P < 0.10\).

DA = divided attention task; PET = positron emission tomography.

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These data are displayed in figure 2. Recognition of shallowly encoded words was tested at EODrcg only. Scores represent median and interquartile range (IQR). Table 5. Recognition of Deeply Encoded Words, in PET Scanner (Rcg1 and Rcg2) and at End of Day (EODrcg)

<table>
<thead>
<tr>
<th>Group</th>
<th>Baseline Shallow Encoding</th>
<th>Baseline Deep Encoding</th>
<th>Drug Shallow Encoding</th>
<th>Drug Deep Encoding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Propofol, n = 6</td>
<td>895 (507)</td>
<td>1,258 (302)</td>
<td>929 (637)</td>
<td>1,346 (101)</td>
</tr>
<tr>
<td>Placebo/DA, n = 5</td>
<td>1,178 (624)</td>
<td>1,684 (438)*</td>
<td>1,196 (404)</td>
<td>1,482 (466)</td>
</tr>
<tr>
<td>Thiopental, n = 6</td>
<td>1,046 (221)</td>
<td>1,367 (182)*</td>
<td>1,207 (383)</td>
<td>1,340 (198)</td>
</tr>
</tbody>
</table>

Data represent correct responses only and are shown as median (interquartile range).

* Reaction time to categorize deep encoding stimuli is greater than shallow encoding (Wilcoxon signed rank, P < 0.05).

DA = divided attention task; PET = positron emission tomography.

(Wilcoxon signed rank, baseline vs. drug encoded words in Rcg2, \( P = 0.028 \), as shown in fig. 2, baseline vs. drug in Rcg2).

To summarize, propofol did not affect reaction time or accuracy during encoding but resulted in memory impairment at the end of the study day. Divided attention and thiopental manipulations likewise did not affect encoding performance but, as opposed to propofol, did not impair memory for deeply encoded words at the end of the study day.

**Regional CBF Correlates of Deep Encoding**

The levels of processing manipulation, requiring semantic categorization, resulted in memories that were recognized at the end of the study day. PET imaging of brain activity at the time of encoding of these memories revealed increased rCBF in a specific region of the left prefrontal cortex when deep encoding was contrasted with shallow encoding in the baseline condition. Because all groups received the same treatment in the baseline condition, image data from all 17 participants were used to define the *a priori* ROI used in subsequent analyses.

The largest region of brain activation during deep encoding was indeed in the left inferior prefrontal cortex as previously reported by other investigators. This ROI (roiPFC) comprised a volume of 239 voxels (6.55 cm², cluster level \( P < 0.001 \) for occurrence of a cluster of voxels this large), with a maximum T value of 7.44 at \( x, y, z \) coordinates in Montreal Neurologic Institute brain space of \(-48, 12, 3\) mm (fig. 1). This region encompassed the middle and inferior frontal gyri, the precentral gyrus, and insula and included Brodmann areas 44-47, 9, 11, and 13.

To address the hypothesis that the amnesic effect of propofol is mediated by an effect that weakens encoding, the response of the roiPFC to deep *versus* shallow encoding was tested separately in each group and condition, *i.e.*, a \( 3 \times 2 \) comparison as shown in figure 3 and table 6. Because of the small numbers in each group, formal contrasts between groups could not be conducted. However, the effect sizes of deep *versus* shallow encoding were used to define the *a priori* ROI used in subsequent analyses.
The LoP manipulation was successful in creating memories for deeply encoded words that lasted over the duration of the study and was a more important influence than the learning effect from repeated presentations of words on the deep encoding task. This was demonstrated by the fact that final recognition memory for deeply encoded words was better than for shallowly encoded words in the thiopental group, even though no learning effect was present in that group. These robust memories were likely a result of greater cognitive processing requirements needed for deep encoding, as revealed by longer reaction times and more error-prone categorizations. Importantly, the LoP manipulation that resulted in these long lasting memories activated a well-defined region in the left prefrontal cortex of the brain was still largely present at a time when behavioral performance on the encoding task was unaffected. To be specific, the amnesic dose of propofol in the current study resulted in memory impairment at the end of the study day but did not impair behavioral measures of encoding performance. Therefore, memory impairment was not as a result of sedation, but rather was representative of drug induced amnesia by propofol. Both thiopental and divided attention manipulations influenced activation in the prefrontal cortex, thus demonstrating the sensitivity of the experimental methods to detect changes in this brain region despite small numbers of participants. Therefore, drug-induced amnesia from propofol did not solely result from impairment of encoding.

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**Discussion**

Using rCBF changes in response to a levels of processing manipulation of word evaluations, the current study examined the influence of propofol on brain processes underlying the encoding of verbal information into long-term episodic memory. At amnesic doses of propofol, encoding activity in the left prefrontal cortex of the brain was still largely present at a time when behavioral performance on the encoding task was unaffected. Therefore, memory impairment was not as a result of sedation, but rather was representative of drug induced amnesia by propofol. Both thio- pelntal and divided attention manipulations influenced activation in the prefrontal cortex, thus demonstrating the sensitivity of the experimental methods to detect changes in this brain region despite small numbers of participants. Therefore, drug-induced amnesia from propofol did not solely result from impairment of encoding.

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The current study revealed that propofol is unlikely to have its major effect on long-term memory by impairing encoding processes. In isolation, the small change in effect size of activation in the LIPFC in the presence of propofol, as seen in the top panels in figure 3, would not be revealing. However, when this result is compared with the substantially decreased effect size by divided attention and thiopental manipulations in the drug condition (right-side panels in fig. 3 and table 6), the different nature of the effect of propofol on LIPFC activity is evident. The divided attention manipulation replicates previous findings that show a decrease in LIPFC activity with changes in attention. In those studies, the change in activity in this brain region was related to changes in long-term memory, which were thought to be mediated via attentional mechanisms impacting so-called executive functions.30,32–34 Subregions of the LIPFC have been closely associated with working memory, which in turn is closely related to encoding processes.12,15,15 Previously, we have demonstrated that thiopental can interfere with encoding via effects on working memory. Therefore, the current study raises the possibility that the LIPFC may be sensitive to changes in attention or to sedation.

As we showed previously and as was replicated in the current study, impairment of memory from propofol occurred some time after encoding of information into long-term memory.19 In comparison with the learning effects present in the placebo/DA group and the baseline response in the propofol group, seen as greater success at Rcg2 than at Rcg1 in figure 2, the effect of propofol on memory seemed to start at some point between 18 and

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**Table 6. Voxels with Peak Activations (MNI coordinates)**

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>Propofol</td>
<td>T = 4.88, P = 0.018</td>
<td>T = 4.00, P = 0.063</td>
</tr>
<tr>
<td></td>
<td>x = -44, y = 4, z = 29</td>
<td>x = -45, y = 27, z = 3</td>
</tr>
<tr>
<td>Placebo/DA</td>
<td>T = 4.41, P = 0.035</td>
<td>T = 2.55, P = 0.398</td>
</tr>
<tr>
<td></td>
<td>x = -45, y = 0, z = 27</td>
<td>x = -45, y = 7, z = 28</td>
</tr>
<tr>
<td>Thiopental</td>
<td>T = 4.52, P = 0.030</td>
<td>T = 2.12, P = 0.579</td>
</tr>
<tr>
<td></td>
<td>x = -48, y = 12, z = 3</td>
<td>x = -36, y = 6, z = 24</td>
</tr>
</tbody>
</table>

Effect size of deep vs. shallow encoding contrasts for increased regional cerebral blood flow in the roIPFC (region of interest, prefrontal cortex) in the left inferior prefrontal cortex shown in figures 1 and 3, as estimated by T values. The maximum voxel significance in the roIPFC, corrected for multiple comparisons in this region using small volume correction in Statistical Parametric Mapping 99, is tabulated, along with the location in Montreal Neurologic Institute (MNI) space of that voxel. The most significant voxel location is indicated in figure 3 by the crosshairs.

DA = divided attention task.
42 min after encoding. This is in accord with the imaging findings in the current study that showed propofol did not have much effect on encoding activity in the LIPFC.

We have previously described decreases in rCBF in the prefrontal cortex with amnesic doses of propofol. At first, these findings seem incongruous with the presence of encoding activity in the LIPFC as seen in the current study. Despite decreases in CBF with anesthetic agents, other investigators have noted that brain activation to stimulation is still present. Therefore, increased rCBF in the LIPFC during deep encoding can be consistent with the decreases in rCBF that we previously reported with propofol. Retention of episodic memories is dependent on a network of brain regions, most notably prefrontal, hippocampal/medial temporal, and parietal regions. The word memory task used in our previous study did not isolate encoding processes, as in the current study, and resulted in increased rCBF in both prefrontal and parietal regions. It is possible that the effect of propofol on memory may have been mediated via actions in the parietal rather than in the prefrontal region. This possibility is consistent with changes in event-related potential measures of recognition memory in the parietal region soon after encoding in the presence of propofol, preliminary results of which we have recently presented. Therefore, the effect of propofol on memory processes and neuroanatomical structures underlying its amnesic actions may be more evident in recognition tasks, particularly those involving parietal regions.

In the current study, there was dissociation between final recognition memory at the end of the study day and LIPFC activity at encoding. In contrast with propofol, there was no change in final recognition memory in the divided attention and thiopental groups, despite the lack of activation in the LIPFC in the drug condition. A number of possibilities can be put forward to explain this observation. In the presence of weaker encoding, as indicated by decreased response in the LIPFC in the thiopental and placebo/DA groups, recognition of words at the end of the study day may have engaged memory processes different from those normally used. For example, familiarity (knowing one has experienced something) may have resulted in recognition instead of recollection (a sense of distinct recall), as occurs in the elderly. The use of alternative recognition processes was not available for the participants who received propofol because any memories were degraded by that time, i.e., there was not even a weak trace available for recognition of most memories. Results from the thiopental group raise the possibility that LIPFC activity may be influenced by sedation. Because nonsedative doses of propofol were used in the current study, little influence on LIPFC activation might have been expected if this had been the case. On the other hand, participants receiving propofol may have compensated for the presence of drug with increased attention during encoding, as has been demonstrated in memory function in the elderly. Such cognitive effort may have counteracted an underlying effect to diminish LIPFC activity.

Activity in the hippocampus rather than the LIPFC may relate more directly to subsequent memory. Despite the fact that we did not find changes in rCBF in the medial temporal lobe in this and in our previous study, animal studies suggest that propofol requires the amygdala to exert its amnesic effect. Therefore, methods that identify the hippocampus/medial temporal lobe, such as the subsequent memory paradigm used in event-related functional magnetic resonance imaging studies, may help to resolve this issue.

There are certain limitations that should be kept in mind when interpreting the findings from the current study. The largest is the unanticipated small accrual. This was problematic in that sufficient power was not present to allow recognition-related brain activity to be imaged, which may have provided valuable insight into the actions of propofol on memory after encoding. Fortunately, this was not the case for encoding activity, because the LoP paradigm was sufficiently robust to allow identification of brain activity in the baseline condition in all groups. The use of multiple recognitions of deeply encoded words, as compared with a single recognition for shallowly encoded words, may have exaggerated differences in memory between these items. Therefore, the final memory effect of propofol is affected by multiple influences beyond those present during encoding, and caution should be exercised in linking memory on final recognition testing and brain activity at encoding. It should be noted that there was some decrease in effect size in the propofol group in the drug condition, although not as large as that present in the control groups. Therefore, the possibility still exists that propofol has some influence, albeit not a major one, on the prefrontal cortex.

In conclusion, these initial observations of brain activity during encoding of auditory words reveal no appreciable effect of amnesic, but not sedative, concentrations of propofol on the left prefrontal cortex. The current study supports findings from our previous studies where encoding of information into long-term memory in the presence of propofol seemed to be normal. As opposed to that which occurs in the elderly, the nature of encoding processes underlying normal encoding performance in the presence of propofol seems largely unaffected. However, the LIPFC region of the prefrontal cortex was affected by a divided attention manipulation or the administration of thiopental. Therefore, this region of the prefrontal cortex may be sensitive to changes in attention or sedation during encoding. In light of the fact that amnesic doses of propofol had little effect on the response of the prefrontal cortex to encoding of verbal information into long-term memory, one can conclude that the production of amnesia at these low doses of propofol does not solely reside in interference with encoding processes, nor does it involve an isolated action on the prefrontal cortex.
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