Investigation of the opioid system in absence seizures with positron emission tomography

Peter A Bartenstein, John S Duncan, Martin C Prevett, Vincent J Cunningham, David R Fish, Anthony K P Jones, Sajinder K Luthra, Guy V Sawle, David J Brooks

Abstract
The neuroanatomical and pathophysiological basis of primary generalised absences is uncertain. Administration of endogenous opioids has been shown to result in absence-like seizures in animal models. Positron emission tomography scans were performed in eight patients with primary generalised epilepsy and eight control subjects. Regional cerebral blood flow was measured interictally with $^{15}$O$_2$, after which a 90 minute dynamic study with the opioid-receptor ligand $^{11}$C-diprenorphine was performed. Serial absences were precipitated by hyperventilation for 10 minutes, starting 30–40 minutes after injection of diprenorphine. Absences, with generalised spike-wave discharges on the EEG, occurred for between 10% and 51% of the provocation period. No individual (normal or patient) had any interictal focal abnormalities of cerebral blood flow. After provocation of serial absence seizures, there was increased diprenorphine elimination from the association cortex, but not from the thalamus, basal ganglia, or cerebellum, compared with control subjects and patients scanned without provocation of absences. It was possible to simulate the observed increased diprenorphine elimination following seizures in cerebral cortex using a two tissue compartment model, with an estimated $15–41\%$ decrease in the specific tracer uptake rate constant ($k_t$). These results suggest that endogenous opioids are released in the association cortex at the time of serial absences, lead to increased receptor occupancy, and may have an important role in the pathophysiology of generalised absences.

Methods
PATIENTS
We studied eight patients (five female, three male) with childhood or juvenile absence epilepsy that had not remitted. Their ages ranged from 20 to 49 years (median 32), median weight 62.5 kg (range: 44–81). Eight male normal volunteers aged between 20 and 41 years (median 34), median weight 72 kg (range: 61–90), were also scanned (table 1). Data on two further patients were not analysed because of movement artefact. All patients were on stable medication, had absence seizures every day and could reliably provoke serial absences by hyperventilation. All patients had normal MRI scans. Five patients (numbers 4–8) were restudied interictally, without provocation of serial absences by hyperventilation.

Permission to perform these studies was granted by the Ethics Committees of the National Hospital for Neurology and Neurosurgery and the Royal Postgraduate Medical School, Hammersmith Hospital, London. Approval to administer radiolabelled gases and ligands was obtained from the Administration of Radioactive Substances Advisory Committee of the United Kingdom (ARSAC). All patients and volunteers gave informed written consent after a full explanation of the procedures.
Table 1  Characteristics of patients and control subjects

<table>
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<th>No</th>
<th>Sex/age (years)</th>
<th>Diagnosis</th>
<th>Duration of epilepsy (years)</th>
<th>Medication (total daily dose, mg)</th>
<th>EEG- % SpW*</th>
<th>EEG- % SpW+</th>
<th>Absence induction period (minutes)</th>
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<td>JAE</td>
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<td>CBZ 800</td>
<td>0-7</td>
<td>41</td>
<td>30-40</td>
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</table>

*Before activation period. †During activation period.

% SpW: Percentage time occupied by absences.
JAE: Juvenile absence epilepsy; CAE: Childhood absence epilepsy; PHT: Phenytoin; ESM: Ethosuximide; VPA: Sodium Valproate; CBZ: Carbamazepine; CLOB: Clobazam.

SCANNING PROCEDURE

Scans were performed on a high resolution PET scanner (CTI 931/12/8, CTI, Knoxville, TN, USA), the performance characteristics of which have been previously described by Spinks et al.20 The final image resolution of this system for 15 simultaneously acquired slices is 8.5 x 8.5 x 7.0 mm (at full width half maximum). Subjects were positioned with the glabla-inion line parallel to the detector-rings so that the intercommisural line (AC-PC line) was in the plane of the directly obtained transaxial images. To minimise movement during scanning, the subject’s head was placed in an individually moulded thermoplastic head support. In addition, continuous direct observation of the alignment of marks put on the skin to projected crossed laser lines, and video subtraction with live reference to a frozen baseline image were performed.

A 22 gauge arterial cannula was inserted into the radial artery after subcutaneous infiltration with 1% bupivacaine. The EEG was monitored continuously in all subjects with an eight channel Siemens Minograph instrument using the 10/20 system of electrode placement. A 10 minute transmission scan was collected using a retractable 68Ga/68Ge ring source. Cerebral blood flow was measured interictally with C18O2 (inhalation for two minutes (6 MBq/ml)) and a three minute scan composed of 20 frames.

Diprenorphine was labelled with 11C in the O-methyl group using 11C-iodomethane.21 After decay of C18O2, 300–500 MBq of 11C-diprenorphine were injected intravenously over 30 s and 26 scans were collected over 90 minutes. The median injected dose of the administered tracer was 353 MBq (range: 312–485) for the patients and 383 MBq (range: 307–449) for the normal subjects. The median specific activity of the tracer was 13 400 MBq/μmol (range: 6100–18 400) for the patients and 11 900 MBq/μmol (range: 6900–22 500) for the normal subjects.

Serial absences were induced by voluntary hyperventilation between 30 and 40 minutes after tracer injection in five patients and between 40 and 50 minutes post-injection in three others. The eight normal volunteers hyperventilated between 30 and 40 minutes post-injection. The arterial pCO2 was measured at 20 minute post-injection and one minute before the end of hyperventilation.

Continuous arterial blood sampling to measure radioactivity was carried out at a rate of 5 ml per minute for the first 10 minutes and then 2.5 ml per minute using an on line bismuth germanate detector system. The formation of diprenorphine metabolites was measured in arterial blood 5, 10, 20 and 45 minutes post-injection.22

DATA ANALYSIS

11C-Diprenorphine scans were analysed with interactive image analysis software (Analyze version 3.0, Biodynamics Research Unit, Mayo Foundation, Rochester, MN, USA) on Sun Workstations (Sun Microsystems Inc, Mountain View, CA, USA). The AC-PC line was defined on an image of the distribution of radioactivity integrated from 30–90 min post-injection.23 In all subjects pitch and roll relative to this line were under three degrees, so reformatting of transaxial scans was not necessary. Images of the distribution of radioactivity integrated for the periods 10–30 minutes and 70–90 minutes were over laid as a further check for subject movement during a scan. Two out of 10 patients scanned showed significant movement (> 5 mm) and were therefore excluded from further analysis. A template of 155 rectangular regions of interest, average size of 1.2 x 1.2 cm, was adapted to the individual brain scans by visual inspection with reference to the stereotactic atlas of Talairach and Tournoux,24 using an integrated image of 11C-diprenorphine related radioactivity collected from 30 to 90 minutes (fig 1). The regions of interest template was placed on the following regions (the number of 7 mm transaxial planes contributing to each region are indicated in parentheses): brainstem (six), thalamus (two), caudate (three), putamen (three), cerebellum (four), anterior (six) and posterior (five) cingulate gyrus, frontal (eight), sensorimotor (five), medial (three) and lateral (five) temporal, parietal (five) and occipital neocortex (three). Time-activity curves, corrected for the physical decay of 11C, were constructed for individual regions of interest and curves of those regions belonging to each of the 13 predefined anatomical regions were combined and averaged.
Analysis was in two parts. First, to control for the differences in injected radioactivity, the possibility of different volumes of tracer distribution in brain tissue, and for the differences in transport of tracer between plasma and cerebral extracellular space in different individuals, the decay-corrected regional time-activity curves were normalised to an amplitude of 100 over the period of 15–30 minutes post-injection. This period was chosen, as opposed to normalising to peak activity, as the time-activity tissue curve had flattened and so was less blood flow dependent, and also because it immediately preceded provocation by hyperventilation. The normalised tracer washout in the period from 60 to 90 minutes post-injection was then compared between all patients and normal subjects with analysis of variance.

Secondly, a computed simulation was performed using the average time-activity curves of the five patients who hyperventilated from 30 to 40 minutes, comparing them with the averaged data of the eight normal subjects, using the primary visual cortex as reference tissue. Estimation of the extent of possible changes in the rate constants (K1, k2, k3 and k4) during the period of hyperventilation was based on the reference tissue model described in detail by Cunningham et al (fig 2). This model assumes that specific binding of the ligand is negligible in a reference region and that the ratio of the rate constants describing the transport of ligand between the vascular and tissue compartments is the same in a reference region \( (K_1/k_2) \) as in a region in which specific binding occurs \( (K_1/k_2) \). A nonlinear least mean squares fit of the tissue time-activity curves from the two regions was carried out on the averaged data of the normal subjects, in order to estimate the binding potential \( (k_2/k_4) \).

The effects of changing the values of \( K_1, k_2, k_3 \) and \( k_4 \) for the duration of the 10 minute provocation period only, and also for the remainder of the study, on the shape of the computer-simulated curve were determined (fig 3), and the results compared with the fit of the tissue time-activity curves from the patients. In this way we sought to estimate the degree of change in the binding potential \( (k_2/k_4) \) at the time of absence seizures and to test the robustness of this result in relation to possible changes in the other compartmental transfer constants. A decay and metabolite corrected arterial input function from a normal subject was used in the computed simulation; \( K_1 = 0.038 \) per minute, \( k_2 = 0.096, k_3 = 0.3, k_4 = 0.06. K_1 \) and \( k_2 \) were calculated from the Renkin-Crone equation:

\[
K_1 = \text{flow} \times [1 - e^{-\frac{1}{\text{flow} \times 30}}], \quad k_2 = \text{flow} \times [1 - e^{-\frac{1}{\text{flow} \times 30}}].
\]

\( PS_{in} = 0.6 \) per minute, \( PS_{out} = 0.15 \) per minute, in which \( PS_{in} \) and \( PS_{out} \) are the apparent permeability-surface area products for transfer of tracer across the blood-brain barrier.

**Results**

In all patients and normal subjects, the interictal \(^{18} \text{O}_2\)-scan showed no focal abnormalities of cerebral perfusion. Serial absences were induced during the 10 minute period of hyperventilation in all patients. Generalised spike-wave discharges occupied between 10% and 51% (median 37%) of the provocation period (table 1; fig 4). The amount of spontaneous spike-wave activity occurring between diprenorphine injection and provocation by hyperventilation was 0.3–7.0% (median 0.7%). The amount of spike-wave activity returned to the baseline rate over 1–5 minutes after the end of hyperventilation. The five patients restudied interictally, without hyperventilation, had spike-wave activity for 0.2–5%, median 1.5%, of the duration of these scans. The EEGs of all control subjects were normal.

The rate of \(^{14} \text{C}\)-diprenorphine metabolism

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**Figure 1** PET image showing regional distribution of \(^{14} \text{C}\)-diprenorphine related radioactivity integrated from 30 to 90 minutes in a normal subject.

**Figure 2** Two tissue compartment model incorporating a reference region and a region in which there are saturable binding sites for \(^{14} \text{C}\)-diprenorphine. \( k_3 = f_k a (B_{max} - B) \); \( k_3 = k_{df} \); binding potential \( = k_2/k_{df} \); \( f_f \) = tissue free tracer fraction.
was similar in patients and normal subjects. Twenty minutes after injection the median percentage of unmetabolised $^{11}$C-diprenorphine in arterial blood was 43.3% (range: 18.6–51.1) in the patients and 43.7% (range: 31.9–57.8) in the normal subjects. The mean fall in arterial pCO$_2$ after hyperventilation was 3.0 kPa (SD 1.1) in patients and 2.8 kPa (SD 0.77) in control subjects.

Analysis of the normalised time activity curves over 60–90 minutes showed a greater clearance of radioactivity following hyperventilation induced serial absences from the brainstem ($p < 0.01$) and association neocortex (frontal, parietal, temporal) ($p = 0.06$), most marked in parietal cortex ($p < 0.01$) and posterior cingulate ($p < 0.05$), compared with normal subjects. There was no increase of tracer washout following serial absences in sensorimotor cortex, occipital cortex, thalamus, caudate, putamen, or cerebellum. There were no side to side differences of diprenorphine binding in regions in which time-activity curves of patients and control subjects did, and did not, diverge. There was no significant difference in time-activity curves before hyperventilation and in all cases the changes were noted after the onset of the flurries of absences (figs 5 and 6).

In the five patients who were studied with and without hyperventilation there was increased tracer washout following the provocation of serial absences from association neocortex (frontal, parietal, and temporal) ($p < 0.01$) (table 2, fig 6), with all areas showing a similar effect. There was no significant difference in tracer washout from the brainstem, or other brain areas, between the scans performed at rest and with provocation of absences. There was no appreciable difference in tracer washout from association cortex in controls and patients studied interictically (fig 6).

The extent of increased tracer washout in patients following hyperventilation was not correlated with the number or duration of absences provoked, the time taken for absences to subside to the baseline rate after the end of hyperventilation, the rate of diprenorphine metabolism, the severity or age of onset of epilepsy, age at time of study, or medication taken.

Comparison of normalised time-activity curves of the control subjects with normative data and simulated time-activity curves showed no increase in the loss of tracer after hyperventilation from any cerebral region. When the averaged time-activity curves of the five patients who hyperventilated from 30 to 40 minutes and the eight normal subjects...
were compared by computed simulation, it was estimated that, if a sustained decrease in $k_1$ underlay the reduced diprenorphine retention seen in the patients, the extent of the fall was $41$ (SD $15\%$) in parietal cortex, $32$ (SD $8.5\%$) in posterior cingulate, $37$ (SD $5.5\%$) in lateral temporal cortex and $15$ (SD $5\%$) in frontal cortex. These results were the consequence of a simulated reduction in $k_2$, from the time of provocation of absences that was continued to the end of the study. If a transient fall in $k_2$, from 30 to 40 minutes only, was simulated, there was less effect on the simulated time-activity curves. The same endpoint, at 90 minutes, was reached by an $80\%$ reduction of $k_2$, from 30 to 40 minutes, and by a $20\%$ reduction of $k_2$, that persisted from 30 to 90 minutes (figs 3b, 3c). The essential parameter that is altered in this simulation is the binding potential $k_1/k_2$; doubling $k_2$ has an equivalent effect to halving $k_1$. Simulation of the effects of changes in cerebral blood flow showed that a four-fold and ten-fold increase in blood flow, from 30 to 40 minutes, had only minimal effects on the diprenorphine time-activity curve (fig 3a).

**Discussion**

The principal finding of this study was an increased loss of $^{11}$C-diprenorphine associated tissue radioactivity following serial absence seizure, from association cortex, compared with patients scanned interictally and hyper-ventilating control subjects. These observations suggest that the reduced tracer retention was associated with the flurry of absences in itself and was not a result of the condition of epilepsy or of hyperventilation. The effect was site selective. Thalamus, basal ganglia, cerebellum, occipital, and sensorimotor cortex did not show increased loss of $^{11}$C-diprenorphine signal after serial absences. Although there was reduced retention of $^{11}$C-diprenorphine in the brainstem of patients after serial absences compared with controls, there was no significant difference between patients studied with and without provocation of absences. In consequence no definite conclusions about changes in diprenorphine binding in the brainstem, in association with serial absences, can be drawn.

The divergence of the ictal and control time-activity curves appeared to follow the occurrence of serial absences, at 30–40 and 40–50 minutes, and there was no evidence to suggest that this preceded the development of absences. The temporal resolution of the dynamic changes in diprenorphine binding was limited, however, and it was not possible to identify definite differences between the time-activity curves of patients who hyper-ventilated at 30–40 or 40–50 minutes. Further, absences occurred in flurries throughout the provocation period and signal to noise limitations precluded a more detailed temporal analysis of the dynamic changes in $^{11}$C-diprenorphine binding during this time.

In the patient group, the extent of increased diprenorphine washout after hyper-
ventilation did not correlate with identified clinical factors (medication, age of onset, duration of epilepsy, occurrence of tonic-clonic seizures), frequency of absences at baseline and on provocation. The basis of the variation in degree of response is uncertain and merits further investigation. This investigation was limited to subjects aged over 18 years by radiation protection regulations. The patients were highly selected, on the basis of having typical absences that had not remitted and which could be precipitated by hyperventilation. In consequence, it is not certain whether the results of this study could be extrapolated to apply to the generality of younger patients with childhood or juvenile absence epilepsy.

Great care was taken to eliminate the confounding effects of subject movement. Continuous EEG monitoring enabled precise quantification of the amount of spike-wave activity in the patients at the time of hyperventilation. No absences occurred during the central blood flow scans. Our data exclude differences in \(^{11}\)C-diprenorphine metabolism between patients and control subjects, and any effects of hyperventilation in themselves, as causes of the peri-ictally increased loss of tissue tracer in the patients. The quantities of the cold diprenorphine injected together with \(^{11}\)C-diprenorphine were always less than 0-3 \(\mu g/kg\), and this has been shown to have a negligible effect on receptor occupancy in vivo.\(^{26}\)

The computed simulation, based on a two tissue compartment model, indicated that even a 10-fold increase in cerebral blood flow at the time of provocation of absences would have only a marginal effect on the tracer kinetics and could not account for the observed \(^{11}\)C-diprenorphine washout during seizure activity. Further, the decreased \(^{11}\)C-diprenorphine binding after serial absences was site selective, whereas the increase in cerebral metabolism and blood flow that has been demonstrated during absences is diffuse and global, including caudate and thalamus.\(^{27}\)

It seems likely that our observation of increased tracer washout after seizures was the consequence of a fall in binding potential for diprenorphine. In the cerebral areas in which peri-ictal changes were observed, the value of \(k_3\) is about four to six times higher than the values of \(k_4\) (fig 2).\(^{18,26}\) A simulation with typical values of \(k_3\) and \(k_4\) showed that doubling \(k_4\) had approximately the same effect on a simulated time-activity curve, as halving \(k_3\). Our data suggest a fall in binding potential \((k_3/k_4)\) following serial absences but do not allow differentiation between changes in \(k_3\), the specific tracer association rate constant, and \(k_4\), the dissociation rate constant. The most likely explanation for our data is a fall in \(k_3\) following serial absences, caused by a decrease in the number of opioid receptors available to bind labelled diprenorphine, rather than an increase in tracer dissociation \((k_3)\). Opioid receptors have a turnover time of the order of days, making it unlikely that changes in total receptor density are the explanation of our data.\(^{29}\) Release of endogenous opioids at the time of, and following absence seizures, which then compete with \(^{11}\)C-diprenorphine for specific binding sites would be expected.

The extent of the observed changes indicates a marked biological effect. Complete arterial input curves were not available for all subjects, so the simulation was based on occipital cortex as a reference input function. The occiput has about 15% of the opioid receptor density of the average neocortex.\(^{18}\) Though this is comparatively small, it results in a slight overestimation of the changes in \(k_3\) in our computed simulation.

The computed simulations provided an estimate of the magnitude of the fall in binding potential that would explain the data. If binding potential was only reduced for the 10 minutes of the provocation period and then returned to baseline values, a much larger reduction of binding potential would be necessary to fit the experimental data than if the reduction of binding potential were maintained over the subsequent 60 minutes of the study (figs 3b, 3c). The gradual divergence of the control and postictal time-activity curves would favour a more modest reduction of binding potential that persists for the remaining 60 minutes of the scan, but our data do not exclude the possibility of a more marked short term effect. Our estimation of 15 to 41% reduction in binding potential is of similar magnitude to the approximately 30% reduction of \(^3\)H-diprenorphine binding seen after depolarisation induced endogenous opioid release in hippocampal slices, and was similar to the 15–30% reduction in \(^{11}\)C-carfentanil binding in amygdala, temporal and cingulate cortex and thalamus found after electroconvulsive treatment and attributed to increased receptor occupation by endogenous opioids.\(^{14,29}\) We suggest therefore that our data indicate the release of endogenous opioids at the time of serial absence seizures in the association areas of cerebral cortex and not in thalamus, basal ganglia, or cerebellum. It is unclear whether diprenorphine binding in the brainstem is affected by serial absences.

The induction of absence-like seizures in rodents by enkephalins and prevention by ethosuximide, valproate and opiate antagonists suggest that enkephalins may have a role in the genesis of absences.\(^{10,11,13}\) In contrast, the release of endogenous opioids, following convulsive seizures, has been suggested as a mechanism for suppressing further ictal activity.\(^{12,20,31}\) The patients we investigated did not have convulsions in the course of their PET studies. Although it is possible that endogenous opioid release may have been an epi-phenomenon to the provocation of serial absences, we speculate that these data indicate release of opioids, in vivo, that have a role in the pathophysiology of the absence seizures. There was no correlation between the number of absences provoked and the extent of reduced diprenorphine retention, suggesting the possibility that opioid release is
one factor in a multifactorial process. The limited temporal resolution of the study does not allow determination of whether opioid release precedes, or is subsequent to, occurrence of absences, as these occurred in flurries over the 10 minute provocation period.

It is of note that we found relative suggestive of endogenous opioid release in the cerebral cortex but not in the thalamus. Thalamocortical pathways are implicated in the generation of absences and it is possible that the effects of endogenous opioids on this circuit are mediated through the association areas of the neocortex.

In a mouse model of generalised absence epilepsy, methionine enkephalin levels were increased in cortex, striatum, and brainstem, with no increase in /-endorphin or dynorphin levels, suggesting involvement of enkephalins in the pathophysiology of generalised absences. It has not been established whether there are any persistent abnormalities of opioid receptor binding in generalised absence epilepsy. Further, the interrelationship between putative endogenous opioid release at the time of seizures and interictal receptor numbers is uncertain at present. The current study has not suggested a difference in cerebral diprenorphine binding between controls and patients, when serial absences are not provoked. Minor differences, however, cannot be excluded and the quantification of regional cerebral diprenorphine binding potential in patients with primary generalised epilepsy, interictally, is the subject of a continuing investigation.

In contrast, in patients with mesial temporal epileptic foci, a 35% up-regulation of opioid binding to /-receptors has been found in overlaying lateral temporal neocortex. In gerbils prone to generalised tonic-clonic seizures, the Bmax (receptor density) of 3H-dihydromorphine binding was increased in substantia nigra, peri-aque ductal grey, and medial geniculate. These data suggest the possibility of tonic up-regulation of opioid neurotransmission that may be related to the pathogenesis of seizures, susceptibility, or reflect homeostatic activation of an endogenous, antiepileptic mechanism at critical sites in subjects with partial and generalised tonic-clonic seizures.

Diprenorphine is a non-selective opioid receptor ligand. Smaller doses of opioid peptides induce seizure activity than have analgesic effects and high doses of naloxone are required to block enkephalin-induced, absence-like seizures, implying involvement of non-/-receptors. Antagonists have been shown to block enkephalin-induced non-convulsive seizures. The relative importance of /-, /, and - receptors, however, in the pathophysiology of different seizure types in vivo remains uncertain. Further studies with more selective probes may identify which receptor subtypes are involved in the pathophysiology of epileptic seizures. In conclusion, this study has produced some evidence for the release of endogenous opioids at the time of serial absence seizures in humans in the association areas of cerebral cortex, that may have an important role in the pathophysiology of absences.

We are grateful to the Medical Research Council, the Deutsche Forschungsgemeinschaft and to the MRC-Cycletron unit staff, especially Professor Richard J B Frackowiak and Dr Tara Jones, for support to C J Simon Shorvon and C P Panayiotopoulos for referring patients.

11 Hoffmann J, Dzoljic MR. Effects of delta opioid antagonists on enkephalin induced seizures. Pharmacology 1987;34:61-5.
Early descriptions of sleep paralysis

Binns is usually credited with the first report in 1842 of paralysis which occurred in a daytime nap; hence his term “daymares”: “utter incapacity for motion or speech, difficult respiration, and extreme dread”.

The report by Binns establishes sleep paralysis in the absence of Gélineau’s narcolepsy; this association was unrecognised until Levin in 1987 identified 16 cases amongst 200 cases of narcolepsy in the literature up to 1933. Weir Mitchell in 1876 noted its occurrence as “nocturnal paralysis” in people who were emotionally and physically healthy.

Macnish in a scholarly account of sleep in 1834, may have depicted sleep paralysis in a nightmare: “At one moment he may have the consciousness of a malignant demon at his side; then to shun the sight of so appalling an object, he will close his eyes, but still the fearful being makes its presence known . . . . if he looks up he beholds horrid eyes glaring upon him and an aspect of hell grinning at him . . . . Or, he may have the idea of a monstrous hag squatted upon his breast — mute, motionless and malignant.” [my italics].

Macnish may have known Henry Fuseli’s celebrated picture (The Nightmare 1781) which shows a demoniacal creature squatting on the chest and belly of a supine woman, apparently trying to rise from her bed; this has been pronounced by Schneck as an example of sleep paralysis.

Kinnier Wilson first introduced the term sleep paralysis for attacks precipitated by a terrifying dream. In Modern Problems in Neurology he described a 26 year old bricklayer with typical narcolepsy (αρχη, numb, torpor) and cataplexy (κατακλύσμα, to strike down). Attacks of “tonelessness” occurred: “Of the greatest interest is the fact that when he has been asleep and dreaming, the emotional content of the dream has precipitated an attack of powerlessness . . . . He was dreaming of a murder . . . . he at once awoke and was fully conscious but was unable to move a single finger . . . . could not make a sound; the more he tried the more intense became his emotion and the more absolute his helplessness; he lay thus, flat on the floor, motionless but suffering acute mental distress, for some fifteen minutes ere the attack dissolved itself spontaneously . . . . Dr MacDonald Critchley, Registrar (sic) . . . . helped materially in the examination . . . . the first neurological examination of a patient in the cataleptic state.”

J M S PEARCE

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