The first electroencephalography (EEG) on a human was reported by Hans Berger in 1920. The fundamental principle of EEG is to record the electrical activity produced by the brain, using electrodes. In considering the role of EEG in psychiatry almost 25 years later, Dennis Williams (1954) observed that studies of EEG recordings had offered the clinical psychiatrist little of positive value. He suggested several reasons for this: EEG does not give a true record of cerebral action potentials; attempts at correlating EEG recordings from millions of cells, the electrical activity of which was not understood, with patients’ clinical picture proved meaningless; the recording techniques used were crude; and EEG records simple physiological changes in neuronal activity which are not reflected in the complex integration of the human brain. Nevertheless, Williams did not dismiss EEG out of hand. He outlined its usefulness in the exclusion of gross organic disorder; the study and modification of local cerebral disorders related to abnormal behaviour; the correlation of changes in EEG recordings with single aspects of temperament or personality; and the use of activation techniques to demonstrate hidden pathologies. There have been technological advances since his erudite article was published, but the fundamental problems remain. In this article, we focus on the role of EEG in current psychiatric practice, EEG findings in specific psychiatric syndromes, clinical indications for requesting EEG, and interpretations of EEG recordings. We will also highlight measures to ensure appropriate and competent use of EEG in the routine clinical practice of psychiatrists.

**EEG in current psychiatric practice**

A retrospective review of EEG requests over a 12-month period found that 6.2% of referrals were made by psychiatrists but that psychiatric referrals had the lowest abnormality detection rate (O’Sullivan 2006). A history of epilepsy, being on clozapine and possible convulsive seizures were found to be the only significant predictors of an abnormal EEG recording. Another study reported the presence of an organic factor identified during clinical assessment to be significant predictor, but none of the abnormal recordings helped in discovering an aetiology that was not already suspected from assessment (Lam 1988). It also reported cases in which psychiatrists ignored abnormal recordings.

Of 187 requests for EEG on patients showing aggressive behaviour, 71% were made to ascertain evidence of epilepsy and 22% to determine the presence of organic brain dysfunction. Only one recording showed unequivocal evidence of an epileptic focus (Stone 2003). The low diagnostic yield reported in the Stone study is similar to the 0.5–2% found in EEG conducted to screen for epilepsy (Gregory 1993). Another issue regarding the diagnostic yield emerged from a study of psychiatric in-patients (Warner 1990): although a third of the EEG recordings were abnormal, this led to a change in diagnosis in only 1.7% of cases.

**Limitations of EEG**

Why is it that EEG produces findings that some consider to be clinically irrelevant? The most important limitation that one should take into account is the disparity in the diagnostic yield of EEG recordings.
account is that, by and large, EEG abnormalities are non-specific. With its poor spatial resolution, EEG cannot pinpoint the pathological process, nor can it aid the differentiation between functional diagnoses. In organic dysfunction, EEG would miss slow-growing, deep-seated tumours. Given that the epileptiform activity in epilepsy is not consistently present, one can expect a false-negative rate of around 50% on interictal EEG recordings (Hopkins 1988) – and a false-positive rate of 2% in the general population (Hughes 1999). Apart from such theoretical and clinical limitations, the cumbersome procedure in itself leads to artefacts that can be difficult to interpret. The comparative lack of psychiatric training among physiologists and the lack of training in observing and interpreting EEG among psychiatrists is another problem.

Despite these shortcomings, EEG is attractive for clinicians: the procedure is non-invasive, can measure spontaneous brain activity, requires a lower level of patient cooperation than functional magnetic resonance imaging, and is relatively inexpensive and easily available.

**The EEG procedure**

An EEG wave is the representation of the differences in electrical potential between an electrode placed on the scalp and a reference electrode placed elsewhere on the head. The transient nature of this electrical difference leads to oscillation, which produces the EEG pattern or trace.

Standard scalp EEG, which uses 21 electrodes, can take up to 40 minutes. In addition to the external placing used in standard EEG, the electrodes can be placed directly inside the skull (intracranial EEG), under the skin of the scalp (subdural) or on the external brain surface and dura mater (electrocochleography).

If abnormality is strongly suspected but the resting EEG is normal, activated EEG is used to increase the probability of recording abnormal patterns. Strenuous hyperventilation and photic stimulation (stimulation of the visual cortex using light) are the most common activation procedures.

In EEG-video monitoring, a standard EEG is conducted with simultaneous video recording of the patient. The advantage of EEG-video monitoring lies in the observation of the temporal relationship between the behaviour under investigation and the EEG recording. Serial and continuous EEG recordings can help to increase the diagnostic yield.

Quantitative EEG (qEEG) utilises analytical algorithms that automatically assess resting and evoked activity of the brain (Box 1). This method involves comparing data from a specific individual with those of database of a large population or of a defined population with a specific disorder.

In qEEG, event-related (evoked) potentials – neuronal electrical responses to stimuli – are extracted from EEG recordings by time domain analysis (analysis of the EEG trace with respect to time). Event-related oscillations are changes in the power spectrum frequency of the ongoing EEG, which can be directly or loosely time-related. They are complementary to the routine EEG as they provide an understanding of higher brain functions, including cognition.

**Physiological factors that influence EEG**

**EEG and age**

Electroencephalogram patterns (Table 1) are influenced by age. Infant EEG recording shows slow-wave and high-amplitude rhythms. The asynchronous, more mature rhythms develop between 2 and 6 years of age. Generalised low-amplitude beta activity starts to appear in puberty, and adult EEG shows posterior alpha and anterior beta activity. In old age, there is a slowing of the alpha frequency accompanied by a decrease in delta activity.

**EEG and state (awake v. sleep pattern)**

Electroencephalograms are also affected by the level of alertness or consciousness. A normal ‘eyes-closed’ wake EEG trace is characterised by highly rhythmic alpha waves (Table 1). Beta waves are not uncommon, especially over the frontocentral regions in a normal adult wake EEG trace. Theta
activity is very limited on wake EEG traces and there should be no delta activity. Alpha coma is a diffuse alpha wave occurring in a comatose state. Table 2 shows the stages of sleep and related EEG patterns.

### EEG in major psychiatric disorders in working-age and older adults

Many associations between specific EEG findings and psychiatric disorders are marred by conflicting reports. In this section we list the robust EEG findings in relation to specific psychiatric disorders and drugs.

#### Delirium

Electroencephalography can be a valuable tool of investigation in delirium. The EEG trace in delirium characteristically shows slowing or drop-out of the posterior dominant rhythm (earliest), generalised theta or delta slow-wave activity, a poorly organised background rhythm, and no reaction to opening and closing of the eyes. Serum anticholinergic activity has been suggested as a biomarker for delirium and such activity has been correlated with the following EEG findings: occipital slowing and qEEG with delta and theta increase as well as alpha decrease, as a result of which there is a lower slow-wave ratio compared with traces obtained in dementia (Thomas 2008a). On qEEG, theta activity and relative power in the delta band collectively distinguish delirium from dementia (Jacobson 1993). Prolonged-activation EEG can be used to diagnose delirium in dementia with 67% sensitivity and 91% specificity based on variations in alpha and delta power density (Thomas 2008b).

#### Dementia

In dementia, qEEG abnormalities include higher frontocentral and parieto-occipital theta values, and lower parieto-occipital beta values. Lower peak frequency has been significantly associated with decline in cognitive function in patients with probable early-stage Alzheimer’s dementia (Clauss 1998). There is evidence that visual and quantitative analyses of EEG traces can be used to differentiate between Alzheimer’s and vascular dementia (Gawel 2009). Quantitative EEG findings indicate that increased EEG slowing is associated with a poorer cognitive performance in patients with vascular dementia (Muresanu 2008).

Electroencephalography can help in discriminating between Alzheimer’s disease and Lewy body dementia at the earliest stages of illness. In a study (Bonanni 2008) that recruited 50 patients with a diagnosis of Alzheimer’s dementia, Lewy body dementia or Parkinson’s disease dementia, dominant wave frequencies were 8.3 Hz (s.d. = 0.6) for the Alzheimer’s disease group and 7.4 Hz (s.d. = 1) for the Lewy body dementia group. Less than half (46%) of the patients with Parkinson’s disease dementia exhibited the EEG abnormalities seen in those with Lewy body dementia.

In a smaller study (n = 19), qEEG results for patients with frontotemporal dementia were marked by no increase in slow activities but by a decrease in fast activities (Lindau 2003). The relative power from the temporal region in the beta-2 band and from the parietal region in the theta, alpha and beta-2 bands are informative qEEG variables in distinguishing between frontotemporal dementia and Alzheimer’s disease, with a diagnostic accuracy of 84.6% (Yener 1996).

### TABLE 1 EEG rhythms

<table>
<thead>
<tr>
<th>Wave type</th>
<th>Frequency, Hz</th>
<th>Feature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delta</td>
<td>&lt;4</td>
<td>Frontal in adults and posterior in children High-amplitude waves Seen in slow-wave sleep of adults</td>
</tr>
<tr>
<td>Theta</td>
<td>4–7</td>
<td>Can be found in young children Transient theta waves can be seen in up to 15% of the normal population</td>
</tr>
<tr>
<td>Alpha</td>
<td>8–12</td>
<td>Bilateral and occipital Higher amplitude on the dominant side Seen in relaxed state with the eyes closed Attenuated with attention</td>
</tr>
<tr>
<td>Beta</td>
<td>12–30</td>
<td>Bilateral, frontal Low-amplitude waves Seen in alert stage</td>
</tr>
<tr>
<td>Mu</td>
<td>7–11</td>
<td>Associated with motor activity and seen in precentral areas Attenuated with contralateral limb movement</td>
</tr>
<tr>
<td>Lambda</td>
<td></td>
<td>Occipital Single sharp wave Associated with ocular movement</td>
</tr>
<tr>
<td>Gamma</td>
<td>34–100</td>
<td>Associated with cognitive and motor functions</td>
</tr>
</tbody>
</table>

### TABLE 2 Sleep and EEG patterns

<table>
<thead>
<tr>
<th>Stage</th>
<th>Feature</th>
<th>EEG pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Transitional stage between sleep and wakefulness</td>
<td>Waves with a frequency 3–7Hz</td>
</tr>
<tr>
<td>2</td>
<td>Fragmentation of thought processes</td>
<td>K-complex and sleep spindles</td>
</tr>
<tr>
<td>3 and 4</td>
<td>Slow-wave sleep</td>
<td>Delta waves of frequency 0.5–1 Hz Stage 4: delta waves more than 50% of the record</td>
</tr>
<tr>
<td>REM</td>
<td>Occurs every 90 minutes</td>
<td>Mixed frequency Saw-tooth waves Low-voltage waves</td>
</tr>
</tbody>
</table>

REM, rapid eye movement.
in dementia and mild cognitive impairment is still insufficient (Jelic 2009).

**Psychotic disorders**

One of the difficulties in interpreting EEG findings that show increased frontal delta activity in people with schizophrenia is that such individuals may be more prone to involuntary saccadic eye movements (Matsue 1986). Antipsychotic medications increase the amount of alpha activity, adding another confounding factor to the EEG recording.

A study of 19 medication-free patients with chronic schizophrenia found greater delta and left-sided beta activity than in the 21 controls (Karson 1988). In the very small number of patients who had reduced alpha frequency, there was an association with larger mean ventricular size. This reduction in alpha frequency has been confirmed in a study of 20 never-treated schizophrenia patients compared with age-matched controls (Omori 1995).

Most research involving qEEG activity power spectra has shown that people with schizophrenia exhibit increased beta and slow-frequency powers and reduced main alpha power (John 1994). One study (Gross 2006), involving 16 patients with treatment-resistant chronic schizophrenia experiencing a relapse, used qEEG to investigate the correlation between the power of different frequency bands over the frontocentral scalp area and Liddle’s factors and negative subscale scores on the Positive and Negative Syndrome Scale (PANSS). The only significant finding was a positive correlation between beta power and psychomotor poverty. Similar studies have shown that negative symptoms are correlated with an increase in beta- and delta-band activity (Williamson 1989), whereas positive symptoms are correlated with increased theta and delta activity (Fehr 2001). Interestingly, a study of qEEG findings in late-onset schizophrenia (n = 10) showed no significant abnormality, thus raising the possibility of a biological difference in the disorder based on the age at onset (Reeves 2003).

**P300 and P50 waves**

The finding of reduction in auditorily evoked P300 wave activity and increased P300 latency among patients with schizophrenia is robust, unlike visually evoked P300 potentials (Box 2). As the visual P300 potentials are influenced by clinical variables, one hypothesis is that auditory P300 is a trait marker of schizophrenia, whereas visual P300 is more of a state marker (Ford 1999). Auditory P300 abnormalities have been detected irrespective of the stage of illness (including presentations in the prodromal stage, in individuals who are actively manifesting symptoms and are considered to be in remission, and in individuals with a family history of schizophrenia) and also in people with schizotypal personality (van der Stelt 2005). These groups also show a prolonged auditory P300 latency. Sagittal reductions in P300 cognitive event-related potentials and reduced left temporal voltage related to asymmetry in the posterior superior temporal gyrus have been reported in patients with schizophrenia. A note of caution is that although the reduction in left temporal voltage has been replicated, many studies have failed to establish it robustly, possibly because of patient selection and response mode.

Abnormal P50 sensory gating correlates due to defects in attention have been found not only in people with schizophrenia but also in their relatives (Clementz 1998).

In a meta-analysis looking at the usefulness of spectral EEG as a diagnostic tool for schizophrenia, Boutros et al (2008) draw the conclusion that most of the studies they looked at confirmed the increased preponderance of slow rhythms in people with schizophrenia, irrespective of their treatment status. However, they point to the lack of robust data on the sensitivity and specificity of the findings and the lack of multicentre studies using standardised criteria. Shagass et al (1984) reported a sensitivity of 50% and a specificity of 90% in a comparison of individuals with schizophrenia and those with depression using slow-wave rhythms as a diagnostic marker.

**Affective disorders**

**Mania**

A review by Shelley et al (2008) reports that EEG abnormalities such as small sharp spikes, 6/s spike and positive spikes are more common in patients

---

**BOX 2 Event-related potentials**

Event-related potentials (ERPs) are recorded while the individual is performing a sensory or cognitive task. They reflect the summated activity of network ensembles active during the task and are characterised by a specific pattern called the waveform, which is composed of negative and positive deflections (waves). For example, a target stimulus detected amid a series of other non-target stimuli produces a positive wave around 300 ms (latency) after the stimulus. This is known as the P300 response. Sensory gating refers to the pre-attentional habituation of responses to repeated exposure to the same sensory stimulus. P50 is an electroencephalogram (EEG) event-related potential waveform used to assess sensory gating.
with mania than in those with depression, in female than in male patients with bipolar disorder, and in non-familial cases of late-age onset mood disorder. Findings such as small sharp spikes, 6/s spike and wave complexes and positive spikes in suicidal patients are of ambiguous clinical significance. There is some evidence to show that interhemispheric asymmetry and decreased coherence of the EEG trace in bipolar disorder can be used to differentiate it from schizophrenia. Anteroposterior P300 topography also differentiates schizophrenia from bipolar disorder with psychotic mania. One study (n = 35 with schizophrenia, n = 20 with psychotic mania) found posterior P300 reductions in the schizophrenia group and anterior reductions in the bipolar group (Salisbury 1999).

Depression
Depression is the most common mental disorder experienced by people with epilepsy (Kanner 2003). Consequently, even if epilepsy has not been diagnosed, EEG findings for people with depression need to be interpreted with caution as they may indicate underlying epileptic disorder rather than being related to depression.

Resting EEG shows greater alpha-wave activity with eyes closed in patients with depression, which has been interpreted as indicating a reduction in cortical activity in depression. Similarly, increased alpha and beta activity has been reported in patients with psychotic depression, but this should be interpreted with caution because such changes can be brought on by metabolic alterations due to poor nutrition (Margerson 1962) or by alterations in adrenaline levels (Gjessing 1967).

Visual P300 (P300 evoked using a visual stimulus) has been found to be abnormal in patients with major depressive disorder and schizophrenia (Blackwood 1987).

Hypnograms showing the pattern of sleep stages through the night reveal irregular transitions between the different sleep states and frequent awakening in people with depression. There is a reduction in deep-sleep stages associated with long periods of rapid eye movement (REM) sleep (Benca 1992). Antidepressants can resolve such irregular transitions and return the EEG pattern during sleep to normal (Figueroa Helland 2008).

Resting EEG has also been used to show that patients with depression who respond to fluoxetine (Bruder 2008), imipramine (Knott 1996), amitriptyline alone and with an adjunctive anticonvulsant or lithium (Ulrich 1986) have a greater alpha power, particularly in the occipital area, than non-responders and healthy controls. Bruder et al (2008) confirmed greater alpha waves (suggestive of lower neuronal activity) over the right than over the left hemisphere, with a high test–retest correlation. Increased pre-treatment activity in the anterior cingulate cortex has been recorded in patients who responded to citalopram and reboxetine when compared with non-responders (Mulert 2007). Deldin & Chiu (2005) report from their EEG study of cognitive restructuring that patients with depression who responded to cognitive treatment exhibited greater overall cortical activity, and cortical asymmetry of greater right relative to left activity in frontal areas, than non-responders.

Pre-treatment alpha measures thus seem to be a predictor of response to antidepressants, but the studies need to be replicated with larger samples to confirm the clinical relevance of these findings.

Anxiety disorders
Although resting EEG recordings from people with anxiety disorders do not show typical deviations from normality, under experimentally induced anxiety, recordings show hypersynchronised alpha activity (Herrmann 1996).

Interestingly, a small study (Arriaga 1991) reported that EEG sleep recordings of participants with primary dysthyemia showed reduced slow-wave sleep and absence of REM sleep disturbances. These indicate that biologically, primary dysthyemia may be closer to generalised anxiety disorder than to affective disorder.

People with panic attacks accompanied by agoraphobic disorders show significant decreases in the power density of the alpha rhythm and increases in the power density of the beta rhythm in the right hemisphere (Gordeev 2008). This reflects a significant activation of the ascending mesencephalic reticular formation. The most characteristic feature in people with panic attacks without agoraphobia is a significant increase in the power density of the theta rhythm in temporal areas of the right hemisphere, reflecting increased activity in temporolimbic structures.

People with obsessive–compulsive disorder (OCD) have increased current density for delta waves in the insula and for beta waves in the frontal, parietal and limbic lobes. Prichep et al (1993) showed that a group of patients who had a relative excess of alpha waves responded well to serotonine reuptake inhibitors for their OCD.

Studies (Jokic-Begic 2003; Chae 2004) show that people with post-traumatic stress disorder (PTSD) have globally reduced complexity in their resting EEG wave forms. However, contrary to almost all hypotheses, Shankman et al (2008) found that individuals with PTSD did not differ from controls on resting EEG asymmetry.
**Personality disorders**

Enhanced cortical slow activity in the delta range has been consistently reported in violent criminal offenders. Increased slow-wave activity in adolescence has been shown to predict emergence of antisocial behaviour later in life (Raine 1990). Attenuated alpha rhythm and increase in theta and delta activity in the frontal lobe has been found in violent offenders with antisocial personality disorder (Reyes 2009). Diminished P300 activity across varied activity has been reported in people with aggression and impulsivity (Gerstle 1998) and in individuals diagnosed with antisocial personality disorder (Bauer 1994).

**Substance misuse and medications**

Alcohol, illicit drugs and prescribed medications (discussed under particular psychiatric disorders) significantly affect EEG recordings. As expected, this influence is greatest on alpha-wave patterns, the dominant pattern seen in the wake state, with increased activity seen in cannabis and cocaine withdrawal and decreased activity in opioid misuse.

In alcohol withdrawal, EEG traces may show attenuation of voltage and prominence of beta activity. The findings of a study (Moeller 1993) that compared the sleep patterns of people with primary major depression and with major depression secondary to alcohol dependence suggest an additive effect of alcohol dependence and comorbid depression on sleep, with changes to delta sleep, REM and non-REM sleep patterns.

**Electroconvulsive therapy**

During electroconvulsive therapy (ECT), EEG recordings show a sequence of high-voltage sharp waves and spikes. This is followed by rhythmic slow waves with a definite endpoint. Wahlund et al (2009) examined frequency distributions of ictal EEG after ECT stimulation in diagnostic subgroups of depression. Psychotic depression has a high occurrence of delta and theta waves, unipolar depression has a high occurrence of delta, theta and gamma waves, and bipolar depression has a high occurrence of gamma waves (accuracy 94%).

**The EEG request**

**When to ask for an EEG**

It is common sense to consider EEG under certain specific circumstances, such as those listed in Box 3. This list is neither prescriptive nor exhaustive, but it implies that clinicians should be highly vigilant for possible organic factors underlying any psychiatric presentation.

**BOX 3 Some situations in which an EEG might be requested**

An EEG might be requested:
- to interpret paroxysmal activity
- to analyse sleep disorders
- to establish the cause of cognitive decline when there is a suspicion of Huntington’s disease, Creutzfeldt–Jakob disease, chronic delirium, etc.

A serial, ambulatory or video EEG may be useful in:
- interpreting periodic behavioural dysfunction
- distinguishing certain types of seizures
- and, to certain extent, differentiating between true and pseudo-seizures

**What information to provide?**

The physiologist or physician who reports on an EEG must have access to all relevant clinical information pertaining to the case being investigated. Box 4 lists the most commonly useful information.

**Training issues**

The most important problem highlighted in studies of current practice is the gap in knowledge among psychiatrists, physicians, physiologists and EEG technicians about utilising and interpreting EEG in patients with psychiatric disorders. Although it is probably practically unfeasible and clinically unnecessary to have an EEG laboratory physically attached to a psychiatric unit, having a laboratory in close proximity to the unit can facilitate collaborative work between psychiatrists and EEG technicians, resulting in a more integrated and appropriate use of EEG (Pogarell 2005).

**BOX 4 Information to submit with the EEG request**

- Description of symptoms, including onset, frequency and duration
- Triggers for those symptoms (e.g. hyperventilation)
- Relevant medical history
- Relevant psychiatric history
- Family history of neurological or neuropsychiatric disorders
- Reports and traces from previous EEGs
- Medications, particularly those known to cause EEG abnormalities
- Results from other investigations
- Differential diagnoses under consideration
Psychiatrists require robust training in the diagnostic usefulness of EEG and interpretation of results, and physicians, physiologists and technicians should have a working knowledge of relevant psychiatric disorders. Psychiatric trainees should be encouraged to participate in case discussions with colleagues regarding EEG. The Royal College of Psychiatrists' curriculum should continue to demand that budding psychiatrists have a good knowledge of this investigation and its clinical relevance. A CPD module on EEG would be of much benefit to both psychiatric trainees and mature clinicians, for whom it might serve as a refresher course.

Conclusions

The application of EEG in psychiatry is in understanding the aetiology and pathological mechanisms of psychiatric disorders, thus aiding clinical diagnosis and guiding appropriate treatment. It can be useful in clinical scenarios where there is a suspected interplay between functional symptoms and organic aetiology. Psychiatrists should base their use of EEG on robust clinical assessment and should be aware of the limitations inherent in the procedure.

References


**MCQs**

Select the single best option for each question stem

1. **Research has shown that:**
   - clozapine therapy is a significant predictor of abnormal EEG recordings
   - the false-positive rate for an epileptiform pattern on EEG traces in general population is less than 1%
   - a third of abnormal EEG traces lead to change in diagnosis
   - psychiatric referrals have a high abnormality detection rate because they are extremely selective
   - the most common request for EEG is to determine the presence of organic brain dysfunction.

2. **In EEG:**
   - auditory stimulation is the most common procedure used in activated EEG
   - the frequency range of recorded waves is between 1 and 20 Hz
   - in old age the trace shows a decrease in delta activity
   - the normal eyes-closed wake EEG trace is characterised by highly rhythmic delta waves
   - standard recording uses 14 electrodes.

3. **The earliest change in EEG due to acute confusional state is:**
   - the appearance of delta waves
   - generalised theta waves
   - intermittent delta activity
   - slowing of the alpha rhythm frequency
   - triphasic waves.

4. **During sleep, the onset of K-complex and spindles is seen in:**
   - stage 1 non-REM
   - stage 2 non-REM
   - stage 3 non-REM
   - stage 4 non-REM
   - REM stage.

5. **In EEG of people with schizophrenia:**
   - increased beta activity is common among patients treated with antipsychotics
   - increased delta activity in the left temporal area is associated with negative symptoms
   - antipsychotic medications decrease alpha activity
   - relatively high mean alpha is a specific finding
   - 6/5 spike and wave complex are common in patients with command hallucinations.
EEG in psychiatric practice: to do or not to do?
Vellingiri Raja Badrakalimuthu, Radhika Swamiraju and Hugo de Waal

APT 2011, 17:114-121.
Access the most recent version at DOI: 10.1192/apt.bp.109.006916

References
This article cites 58 articles, 6 of which you can access for free at:
http://apt.rcpsych.org/content/17/2/114#BIBL

Reprints/permissions
To obtain reprints or permission to reproduce material from this paper, please write to permissions@rcpsych.ac.uk

You can respond to this article at
/letters/submit/aptrcpsych;17/2/114

Downloaded from
http://apt.rcpsych.org/ on October 18, 2017
Published by The Royal College of Psychiatrists

To subscribe to BJPsych Advances go to:
http://apt.rcpsych.org/site/subscriptions/