Epilepsy is common and serious (prevalence 750 per 100 000) and has an impact upon employment, education, and driving. The diagnosis requires a detailed history including witness account. Clinicians must distinguish seizures particularly from syncope and psychogenic attacks. Electroencephalography and magnetic resonance brain scanning help to identify causes and classification of epilepsy, but alone rarely provide the diagnosis. Antiepileptic drug treatment is required long term and is potentially hazardous; patients should start treatment only after informed discussion with an epilepsy specialist. Patients require reliable written information, particularly the driving regulations, and the impact of seizures on employment, education, and leisure. Women must understand the potential drug teratogenic effects. Certain patient groups benefit from targeted epilepsy services, for example, learning disabled, children, teenagers, and elderly. People with epilepsy require long term specialist follow up. Although this is currently provided in mainly in secondary care (including nurse led clinics), improved liaison with primary care should enable improved access to epilepsy services. Epilepsy care should be multidisciplinary and long term, linking primary and secondary care, and empowering patients towards improved management of their condition.

Epileptology has changed from a “Cinderella” specialty to arguably the most exciting area of neurology. We now have a range of effective antiepileptic drugs, high quality imaging identifying a structural basis for most adult onset epilepsy, and an increasing public awareness of epilepsy and its problems. What is still lacking is the number of specialists needed to deliver an essentially clinical and supportive service for a long term disorder affecting almost 1% of the population. Guidance from the National Institute for Health and Clinical Excellence (NICE) on the diagnosis and management of the epilepsies offers national standards for epilepsy care and will help to highlight and correct current deficiencies in provision of epilepsy care.

Epilepsy is the commonest serious neurological condition after stroke (prevalence 750 per 100 000); the workload of its management is potentially huge. However, most patients become seizure free with antiepileptic drugs and can live normal lives, but 30% have continued seizures or significant drug side effects, or both; these require regular epilepsy clinic review. Epilepsy incidence is 50 per 100 000 people per year. However, for each person diagnosed with epilepsy, four to five people with blackouts must be assessed; thus 250 per 100 000 people require specialist assessment.

Three patient groups attend epilepsy clinics:

- Patients for review, usually but not always with epilepsy.
- New patients with undiagnosed blackouts (seizure, syncope, or psychogenic), who need urgent (within two weeks) specialist assessment.
- New patients with an epilepsy diagnosis (not necessarily correct) with particular issues, who need detailed reappraisal.

HISTORY

Diagnosing episodic changes in consciousness requires a general medical perspective, an understanding of the differential diagnosis, and knowledge of seizure and epilepsy classification. A suitably experienced clinician must take a detailed history, including a witness account. There is no short cut, making blackout diagnosis a time consuming activity. Investigations such as electroencephalogram (EEG) and magnetic resonance (MR) brain scanning can support the clinical diagnosis, but generally the history is crucial. Where there is doubt (this is common), re-taking the history is more helpful than repeating tests.

Key components

The history should focus on precipitants (situation and trigger), warning (prodrome), the episode, and the symptoms that follow (recovery).

Table 1 lists characteristics helpful in distinguishing the common causes of blackouts: syncope, seizures, and psychogenic episodes; other transient episodes include migraine, transient ischaemic attacks, and movement disorders.2 The clinician must next decide whether the seizure was provoked or unprovoked, the first (single) seizure or part of a recurrent tendency.

Abbreviations: AED, antiepileptic drug; EEG, electroencephalogram; CT, computed tomography; MRI, magnetic resonance imaging
(epilepsy), and its classification (focal or generalised, idiopathic or symptomatic).

**Provoked or unprovoked?**

Provoked (acute symptomatic) seizures occur with transient cerebral insults. Examples include alcohol withdrawal, drug intoxication, meningitis/encephalitis, head injury, and intracerebral haemorrhage. Long term antiepileptic drug treatment is usually not needed for these.

**Single seizure or epilepsy?**

Most people with epilepsy are diagnosed after a major seizure, but often have had preceding minor events. People may not consider myoclonic jerks, absence seizures, simple or even complex partial seizures to have been epileptic events, and so they go unreported.

**Classification**

Seizures are either generalised or focal.

- Primarily generalised seizures include typical absences (abrupt onset and offset, 3 Hz spike and wave on EEG, usually normal intellect), myoclonic jerks, and generalised tonic-clonic seizures.
- Focal (partial onset) seizures include déjà vu or epigastric aura of medial temporal lobe epilepsy, head and eye turning (adverse seizure) of frontal lobe epilepsy, or visual aura of occipital epilepsy. Generalised tonic-clonic seizures in adults are usually secondarily generalised.

Epilepsies are classified as generalised or focal (localisation related) according to the predominant seizure type, but also takes note of the possible underlying cause:

- Idiopathic epilepsies typically have age specific onset (child or adolescent), favourable response to antiepileptic drugs (AEDs), normal cerebral imaging, and a presumed genetic aetiology. Idiopathic generalised epilepsies (for example, juvenile myoclonic epilepsy) comprise 30% of epilepsies and present with combinations of generalised seizures.
- Symptomatic epilepsies have a known underlying cause (usually structural), such as mesial temporal sclerosis (fig 1), tumour or cortical dysplasia. They mainly have focal seizures, often resistant to drug treatment. Cryptogenic epilepsies have a presumed symptomatic cause, but a definite explanation cannot be found (usually normal imaging).

**Drug treatment history**

This should include:

- Current and previous AEDs, including dose, formulation, dates, benefit, and adverse effects.
- Potentially epileptogenic drugs, for example, ciprofloxacin, tramadol, antimalarials.
- Drugs with important AED interactions, for example, warfarin, digoxin, oral contraceptive.

**Medical history**

Previous blackout events must be explored in detail and patients asked specifically for any history of absences, myoclonus, or photosensitivity. The history should include potential cerebral insults, for example, premature and/or traumatic birth, febrile seizures, meningitis/encephalitis, and

<table>
<thead>
<tr>
<th>Syncope</th>
<th>Seizure</th>
<th>Psychogenic episode</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trigger</td>
<td>Common (upright, bathroom, blood)</td>
<td>Rare (flashing lights, hyperventilation)</td>
</tr>
<tr>
<td>Prodrome</td>
<td>Almost always</td>
<td>Common (aura)</td>
</tr>
<tr>
<td>Onset</td>
<td>Gradual</td>
<td>Usually sudden</td>
</tr>
<tr>
<td>Duration</td>
<td>1–30 seconds</td>
<td>1–3 minutes</td>
</tr>
<tr>
<td>Colour</td>
<td>Very pale</td>
<td>Cyanosed</td>
</tr>
<tr>
<td>Convulsions</td>
<td>Common (brief)</td>
<td>Common (prolonged)</td>
</tr>
<tr>
<td>Eyes closed</td>
<td>Often</td>
<td>Less common</td>
</tr>
<tr>
<td>Incontinence</td>
<td>Uncommon</td>
<td>Common</td>
</tr>
<tr>
<td>Lateral tongue bite</td>
<td>Rare</td>
<td>Common</td>
</tr>
<tr>
<td>Breathing</td>
<td>Quiet</td>
<td>Apnoea (expiration)</td>
</tr>
<tr>
<td>Post-ictal confusion</td>
<td>Rare</td>
<td>Common</td>
</tr>
<tr>
<td>Recovery</td>
<td>Rapid (wakes on floor)</td>
<td>Slow (wakes in ambulance)</td>
</tr>
<tr>
<td>Self injury</td>
<td>Rare</td>
<td>Common</td>
</tr>
</tbody>
</table>

**Table 1** History points distinguishing syncope, seizures, and psychogenic episodes. Note the syncope features here relate to vasovagal syncope: cardiac syncope (even when tachycardia related) may occur abruptly and without prodrome.

**Figure 1** Schematic brain section illustrating left sided mesial temporal sclerosis. Note asymmetry of hippocampi, temporal lobes/cortex, and fornices.
head injury. Heart disease (congenital or acquired) may suggest syncope. Depression and anxiety commonly accompany epilepsy, but significant psychiatric history (including misuse and illicit drug dependence) might favour psychogenic seizures.

**Family history**
The family history should include epilepsy, febrile seizures, syncope, and sudden unexpected death. Family histories are notoriously unreliable, incomplete, sometimes deliberately concealed, and may require repeated inquiry or even direct assessment of affected people.

**Social history**
This includes education, employment, driving status, family planning, home situation, sporting interests, use of alcohol and illicit drugs.

**EXAMINATION**
Physical examination contributes surprisingly little to black-out diagnosis.

- **Epilepsy:** examination includes a search for skin stigmata (neurofibromatosis, tuberous sclerosis), dysmorphic features, body size asymmetry (for example, nail size), and cerebral bruit. Long term AEDs may result in tremor, hair loss, weight gain (for example, sodium valproate, gabapentin), gum hypertrophy, hirsutism, acne, ataxia, or absent reflexes (for example, phenytoin). Patients with focal onset seizures require examination for visual field defects or long tract signs. Field defects from vigabatrin therapy or temporal lobe epilepsy surgery may have implications for driving, even when seizure free.
- **Probable syncope:** cardiovascular examination is essential, particularly in the elderly patient.
- **Psychogenic episodes:** 25% of patients with unexplained blackouts have panic disorder and hyperventilation. It can be helpful (to patient and clinician) to provoke the physical symptoms of hyperventilation in the clinic by deep breathing for three minutes. Hyperventilation may also induce typical absences in children. It is worth looking for wrist scars (previous self harm) and needle marks as predictors of psychogenic episodes.

**INVESTIGATIONS**
All first seizures must be explained and usually investigated. “Everyone is allowed one seizure” is nonsense and potentially dangerous. Note however that normal EEG and brain scan does not exclude epilepsy.

EEG can help to distinguish generalised from focal epilepsies (fig 2), support an epilepsy syndrome diagnosis, and localise the focus of partial seizures. However, it is normal in about 60% of people after a single seizure and in about 40% with epilepsy. EEGs are at their most useful soon after the seizure and before AEDs are prescribed. EEGs are used to support clinical suspicion rather than as the sole means of making the diagnosis of epilepsy. Uncritical interpretation of EEG by those unaware of its limitations presents dangers of erroneous diagnosis of epilepsy, unnecessary restrictions, stigma, and long term treatment. Prolonged video EEG however may capture typical episodes and is particularly helpful in distinguishing epileptic from non-epileptic episodes.

Brain imaging is indicated for spontaneous seizures either of presumed focal onset (aura, focal signs, or EEG focus) or refractory to medical treatment. Spontaneous seizures arising in adults are mostly focal and so all should be considered for cerebral imaging. MRI is the modality of choice because computed tomography (CT) often misses epilepsy causes such as mesial temporal sclerosis (fig 1), cortical dysplasia, cavernoma, and benign temporal lobe tumours, for example, ganglioglioma, dysembryoplastic neuroepithelial tumour.

Electrocardiogram (ECG): 12 lead ECG is indicated after all undiagnosed blackouts, especially suspected syncope and in the elderly patient. Rare cardiac causes of syncope (long QT, Brugada syndrome) mimic epilepsy and can induce sudden death. Investigation of suspected syncope follows standard guidelines, and includes ECG, exercise testing, head up tilt table testing, and 24 hour ECG. Suspected cardiogenic syncope requires urgent cardiology referral.

**MANAGEMENT**

**Starting drug treatment**
AEDs are considered usually after more than one spontaneous epileptic seizure. The MESS study, compared immediate with delayed treatment after single seizures and early epilepsy, and showed that 14 patients were randomised to treatment to prevent one patient relapsing by two years. Long term drug treatment is therefore usually withheld after a single seizure. However, highly epileptogenic causes such as glialoma, justify AEDs after a single event.

AED treatment is usually long term, requiring informed discussion with an epilepsy specialist. Short term AED trials are rarely justified. The patient must balance the seizure morbidity (including the small risk of sudden unexpected death (SUDEP) against the consequences, inconvenience, and the adverse effects for that person (including potential teratogenicity).

**Choosing AEDs**
The 2004 England and Wales NICE guidelines on the use of new AEDs advise initially either carbamazepine (focal seizures) or sodium valproate (focal or generalised seizures).

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*Figure 2* EEGs showing (A) generalised spike and wave activity and (B) left temporal epileptic focus. On each tracing right sided leads are the upper four tracings; left sided leads the lower four.
Women of child-bearing potential (including girls requiring AED during their childbearing years) should probably not be prescribed valproate as first line treatment because of teratogenicity. Lamotrigine seems a safer alternative. Alternative monotherapy should be tried before considering polytherapy. Vigabatrin is no longer started (except in babies with West syndrome) because of problems with permanent visual field constriction. The role of some of the newer drugs (gabapentin, levetiracetam, oxcarbazepine, pregabalin, tiagabine, topiramate) will become clearer after SANAD, a large randomised controlled study comparing first line monotherapy with new and conventional AEDs.

Stopping AEDs
Seizure free patients require detailed discussion before stopping AEDs. In children it is usual to try after two years seizure free. In adults, continued seizure freedom for driving and employment often justifies the inconvenience of continued drug treatment; many adults therefore continue taking AEDs while seizure free for years. Women wishing to conceive naturally want to stop AEDs (see below), and often do so without consulting their doctor. Overall, 40% of adults seizure free for two years will relapse. The risk is highest with previous tonic-clonic or myoclonic seizures, seizures after starting AEDs, needing more than one AED, and in those with abnormal EEGs. The greater likelihood of seizures in the months after withdrawal is reflected in the UK Driver and Vehicle Licensing Agency’s (DVLA) advice to stop driving from the start of AED withdrawal until six months after its completion.

Surgery
Symptomatic epilepsies, for example, from mesial temporal sclerosis, are commonly resistant to AEDs and justify consideration of surgery. Epilepsy surgery is generally underused in the UK. The detailed preparation for surgery (prolonged and sometimes invasive video EEG monitoring, sodium amytal testing) is available in only a few centres. Potentially curative procedures include removal of confirmed epileptogenic lesions including temporal lobectomy for mesial temporal sclerosis; palliative procedures include multiple subpial transection, corpus callosumy, and hemispherectomy in patients with severe symptomatic epilepsies. Vagus nerve stimulation is an option for adults and children with resistant epilepsy.

SPECIAL SITUATIONS
Refractory epilepsy
Refractory focal epilepsy demands a detailed search for structural abnormality. Not all MR scans are equal in their quality of data acquisition or reporting (ideally brain MRI should include high definition, thin sliced images, with FLAIR sequences, and reported by a neuroradiologist); repeat imaging may be necessary. Poor treatment compliance may cause apparent refractory epilepsy. Furthermore, 15% of patients with “refractory epilepsy” have only psychogenic seizures. Patients who have not responded to two first line AEDs therefore require careful diagnostic review. The epilepsy classification must also be reviewed as certain AEDs beneficial to focal epilepsies (for example, carbamazepine, gabapentin) may be ineffective, and even worsened underuse in the UK. The detailed preparation for surgery months after its completion.

Syndrome presenting in old age requires urgent cardiological evaluation. Cerebrovascular disease is closely linked to elderly onset epilepsy; such patients require consideration of antiplatelet and statin treatment, as well as AED. There is little evidence to support any particular AED in the elderly patient; in general AEDs with renal excretion and fewer interactions are preferred.

Pregnancy
Young women need clear information upon which to base treatment and lifestyle decisions (see below). However, for women already pregnant, teratogenicity advice comes too late. Epilepsy nurse input into antenatal clinics helps management of epilepsy in pregnancy (about 0.5% of all pregnancies). Combined neurologist and obstetrician clinics are useful for complicated cases.

Syncope
Syncope affects 22% of the population; and presents potentially an enormous problem. Although only a few are referred to specialists (mainly cardiologists), syncope is still the commonest diagnosis among new referrals to an epilepsy clinic. Epilepsy specialists must therefore work closely with cardiologists, preferably in a joint “blackout” clinic.

Psychogenic episodes
The scarcity of local liaison psychiatry services means that patients with psychogenic seizures are often followed up in epilepsy clinics, becoming major users of epilepsy services. Ideally, one specialist should supervise their management (including sustained AED withdrawal) using regular short interval follow up and admissions only under that specialist’s care. Without this, patients risk admission as “known epileptic” and their AED restarted or increased.

INFORMATION FOR PATIENTS
Driving
Loss of driving privileges contributes significantly to the social predicament of epilepsy. Many people are told that they cannot drive until seeing the specialist, and eagerly await advice. After an unprovoked epileptic seizure (or undiagnosed blackout), UK drivers must stop, inform the DVLA, and remain seizure free for a year before regaining their licence. This law applies even to minor seizures including epileptic myoclonic jerks. Provoked seizures, for example, within a week of head injury, are dealt with individually by the DVLA. For heavy goods and public service licences, drivers must be seizure free for 10 years and off drug treatment.

Lifestyle
People with epilepsy should be encouraged to live normal lives, within sensible limits. Sports and leisure: the seizure frequency and type influence advice for specific circumstances such as
swimming, cycling on busy roads, and isolation sports (for example, horse riding, hill walking, etc.).

Alcohol may provoke seizures through sleep loss, AED interaction (long term alcohol intake induces liver enzymes), forgetting to take AEDs, or inducing misplaced confidence that AEDs can be omitted. Pragmatic advice is that patients with epilepsy should limit alcohol consumption to four units in 24 hours.

Sleep deprivation is an avoidable cause of lowered seizure threshold, especially in idiopathic generalised epilepsies.

Flashing lights: true photosensitivity is uncommon in adults especially while taking AEDs, but many people with epilepsy misguidedly avoid computers, televisions, and discs.

Teratogenicity

Women contemplating pregnancy require balanced and reliable information about the teratogenic potential of their AED. Unfortunately, such data are currently lacking and advice is based to an extent upon opinion and conjecture. Nevertheless, prospective observational data from the UK Epilepsy and Pregnancy Register show valproate to be associated with major congenital malformations more than either carbamazepine or lamotrigine. The risks, although multifactorial, relate to AED burden: monotherapy 4%–6%, duotherapy 7%–8%, and polytherapy 15%–20%. Furthermore, there are suggestions, awaiting prospective evaluation, of increased neuro-developmental delay among children exposed to AEDs in utero. Unfortunately valproate is the AED of first choice for idiopathic generalised epilepsies and so changing drugs to protect unborn children risks compromising seizure control. Also, switching from valproate to lamotrigine is complicated, taking several months. Despite the absence of conclusive proof and the inherent difficulties in researching this area, young women, particularly taking valproate, require specialist review to inform decisions about long term treatment.

FOLLOW UP

Epilepsy, more than many chronic disorders, justifies long term follow up. The diagnosis is history based and too often is made incorrectly, particularly in non-specialist hands. The choice and need for prescribed AEDs may be inappropriate, and patients may too easily accept drug side effects and unnecessary lifestyle restrictions. Good practice would suggest annual review, including in “nurse led” epilepsy specialist clinics, be offered to all patients with epilepsy. Proactive specialist review of those currently managed in the community may also be justified to check diagnoses, optimise clinical management, and to provide information.

AND FINALLY...

Patients need opportunities to ask questions. “What would you like to ask?” induces more response than “Do you have any questions?” Early follow up after the initial diagnosis provides the opportunity to discuss concerns after a period of reflection. Written information is also important, including information sheets, details of web sites, and local support groups. Most powerful is a personalised letter summing up the consultation and giving specific information. Copying the standard clinic letter is a useful alternative.

Patients with chronic conditions are expert in their individual condition as they constantly live with it. Providing individualised verbal and written information and offering specialist nurse telephone contact encourages greater patient involvement in their long term management.

CONCLUSION

Epilepsy clinics provide a multidisciplinary focus for the diagnosis and long term management of patients with recurrent blackouts and epilepsy. The diagnosis is clinical rather than investigation based, and the emphasis of management is long term, team delivered, and founded upon a partnership of specialist care with the patient and with primary care, sharing information, supporting and empowering patients, and aiming for increasing independence.

Authors’ affiliations

S Hadjikoutis, P E M Smith, The Epilepsy Unit, University Hospital of Wales, Cardiff, UK

Conflicts of interest: PS has received hospitality and support from all of the major pharmaceutical companies manufacturing antiepileptic drugs available in the UK.

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