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Scan this QR code for further information around what is epilepsy?

For further information on what is epilepsy visit epilepsysociety.org.uk/what-epilepsy

For a printed copy, or for a large print version, call our helpline on 01494 601 400.
An introduction

Epilepsy varies greatly and affects everyone differently. If you or someone you know has epilepsy, you may find it helpful to learn more. This leaflet gives some basic facts about epilepsy, and points out where you can get more detailed information and help.

Epilepsy

Epilepsy is a neurological condition where there is a tendency to have seizures that start in the brain.

Not all seizures are due to epilepsy. Other conditions that can look like epilepsy include fainting, or very low blood sugar, which may happen in people being treated for diabetes.

In this leaflet when we use the term ‘seizure’ we mean epileptic seizure.

How seizures start

The brain has millions of nerve cells which control the way we think, move and feel. The brain uses electrical signals to send messages from one nerve cell to another. If the messages are interrupted, or the electrical signals do not switch off when they are no longer needed, this can cause a brief change in the way the brain works. This interruption or build up of electrical signals can cause a seizure (sometimes called a ‘fit’ or ‘attack’).
Epilepsy is common

Anyone can develop epilepsy, at any time of life. It happens in people of all ages. There are over half a million people with epilepsy in the UK, so around 1 in 100 people.

There are many different ‘epilepsies’

Epilepsy is not just one condition, but a group of many different ‘epilepsies’ with one thing in common: a tendency to have seizures which start in the brain. Just knowing that a person ‘has epilepsy’ does not tell you much about their epilepsy, or the type of seizures they have.

However, in this leaflet we use the term ‘epilepsy’ as it is a familiar term for many people.

How epilepsy is described

You may see epilepsy described in two ways. The type of epilepsy describes what has caused the seizures to start, and which part of the brain is affected during a seizure. For example, in the term ‘genetic generalised epilepsy’, ‘genetic’ refers to the likely cause (see page 5), and ‘generalised’ means that both sides of the brain are affected during a seizure.

Another way to describe epilepsy is to talk about the type of seizures a person has. In this leaflet we look at the types of epilepsy and not at the types of seizures.

Visit epilepsysociety.org.uk/epileptic-seizures
What causes epilepsy?

Different epilepsies are due to many different underlying causes. The causes can be complex, and sometimes hard to identify. A person might start having seizures because they have one or more of the following:

- A genetic tendency, passed down from one or both parents (inherited).
- A genetic tendency that is not inherited, but is a new change in the person’s genes.
- A structural (sometimes called ‘symptomatic’) change in the brain, such as the brain not developing properly, or damage caused by a brain injury, infections like meningitis, a stroke, or a tumour. A brain scan, such as Magnetic Resonance Imaging (MRI), may show this.
- Structural changes due to genetic conditions such as tuberous sclerosis or neurofibromatosis, which can cause growths affecting the brain.

Some researchers now believe that the chance of developing epilepsy is probably always genetic to some extent, in that any person who starts having seizures has always had some level of genetic likelihood to do so. This level can range from high to low and anywhere in between.

Even if seizures start after a brain injury or other structural change, this may be due to both the structural change and the person’s genetic tendency to have seizures, combined. This makes sense if we consider that many people might have a similar brain injury, but not all of them develop epilepsy afterwards.
Seizure thresholds

Part of the genetic likelihood of developing seizures is called a seizure threshold. This is our individual level of resistance to seizures. Any of us could have a seizure under certain circumstances but, for most people, their natural resistance to having seizures is high enough to stop that happening.

Our seizure threshold is one part of our genetic makeup which can be passed from parent to child. So the chance of you having seizures may depend partly on whether either or both of your parents has epilepsy.

If you have a low seizure threshold, your brain is less resistant to seizures. So you are more likely to suddenly start having seizures for no obvious reason than someone with a high seizure threshold.

Your doctors may be able to tell you what has caused your seizures to start, but this is not always possible. Research continues into understanding more about why seizures happen in some people and not in others.

How is epilepsy diagnosed?

Diagnosing epilepsy can be difficult. Unless someone is having a seizure, there is often no obvious sign that they have epilepsy.

Many people will have a one-off seizure at some point in their lives, but a diagnosis of epilepsy is usually made after a person has had more than one seizure.
The person who has the seizure may not remember what happened. So it can be very helpful to have a description of what happened, from someone who saw the seizure, to pass on to the specialist.

A number of investigations may give more detailed information that can help with a diagnosis. These include:

• blood tests;

• an electroencephalogram (EEG), a test where electrodes are used to record brain activity;

• a brain scan called Computerised Tomography (CT scan), which is a special type of x-ray using a scanner and a computer to take pictures of the brain; and

• Magnetic Resonance Imaging (MRI). This is a type of scan that uses strong magnetic fields and radio waves to produce detailed pictures of the brain.

However, these tests alone cannot confirm or rule out a diagnosis of epilepsy. Often it is a combination of test results, a person’s medical history, and information from those who saw the seizure, that is used to reach a diagnosis.

Visit epilepsysociety.org.uk/diagnosis
How is epilepsy treated?

Anti-seizure medication (ASM)

Up to 70% of people with epilepsy could have their seizures controlled (they stop having seizures), with the right medication. ASM is taken regularly to prevent seizures from happening, by reducing the brain’s excessive electrical activity. ASM is not used during a seizure to stop it, and it does not cure epilepsy. There are different ASMs, and the ASM that someone is prescribed depends partly on the type of seizures they have.

Some people’s epilepsy goes into spontaneous remission and they stop having seizures. When this happens they may be able to gradually stop taking their ASM, with guidance from their neurologist. Other people need to carry on taking ASM for the long term to keep their seizures under control. This may be the case if the underlying cause of the seizures is still there, for example a scar on the brain.

Other treatments for epilepsy

For some people whose epilepsy does not respond to ASM, other treatment options may be possible. These include the following:

- Epilepsy surgery (also called neurosurgery) may be possible for some people if tests are able to show where in the brain the seizures are starting.
• VNS (Vagus nerve stimulation) therapy aims to help regulate the brain’s normal electrical activity to reduce the number or severity of seizures. It is a form of treatment that sends mild electrical stimulation to the brain through the vagus nerve (a large nerve in the neck). It is normally used alongside ASM.

• For some children and adults, the ketogenic diet may help to reduce the number or severity of their seizures. The diet is a medical treatment, often started alongside ASM, and is supervised by trained medical specialists and dietitians. Dietary treatments for adults are available on a limited basis in the UK. Visit epilepsysociety.org.uk/epilepsy-surgery epilepsysociety.org.uk/vagus-nerve-stimulation epilepsysociety.org.uk/ketogenic-diet

Seizure triggers

For some people, certain situations can trigger (set off) a seizure. Common triggers include lack of sleep, stress, alcohol, and not taking their prescribed ASM. Less commonly, seizures can be triggered by flashing lights or moving patterns (called photosensitive epilepsy). It may be possible to avoid triggers if they can be identified.

Visit epilepsysociety.org.uk/photosensitive-epilepsy
**Will epilepsy affect my life?**

Developing epilepsy may affect you in different ways. Some people may feel relieved to be given a name and treatment for their condition.

Sometimes a diagnosis can be hard to come to terms with. Talking about any worries you may have, asking questions, and sharing information may help you, and your family and friends, to make sense of what is happening for you.

Visit epilepsysociety.org.uk/diagnosis

Some people find that their epilepsy does not affect their life much, especially if their seizures are controlled with treatment that suits them. For other people, epilepsy may affect different areas of life, such as work or home life, lifestyle, travel and holidays, leisure, or social life.

Visit epilepsysociety.org.uk/work-employment-and-epilepsy
Visit epilepsysociety.org.uk/travel-and-holidays
Visit epilepsysociety.org.uk/sports-and-leisure
Visit epilepsysociety.org.uk/wellbeing

However epilepsy affects you, or how you feel about it, there is help available.

Visit epilepsysociety.org.uk/what-help-available

There can be a lot of information about the condition to come to terms with. It is important to remember that epilepsy is a very individual condition and affects people differently.
Support available

Epilepsy Society helpline

Our confidential helpline provides emotional support and information for anyone wanting to know more about epilepsy.

Call our helpline (see back cover).

How to help

Epilepsy can happen to anyone. Would you know what to do if you saw someone having a seizure?

Remember the three ‘C’s:

1. Calm - stay calm and take control of the situation.
2. Cushion - cushion their head with something soft.
3. Call - call an ambulance.

Visit epilepsysociety.org.uk/ccc or for more information visit epilepsysociety.org.uk/seizure-first-aid

Every effort is made to ensure that all information is correct at the time of printing. Please note that information is intended for a UK audience. This information is not a substitute for advice from your own doctors. Epilepsy Society is not responsible for any actions taken as a result of using this information.
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