For further information and details of helplines for people with epilepsy, please visit sudep.org or call 01235 772850.
As someone with epilepsy, you’ve almost certainly had to think about risks. Simple precautions – such as taking a shower rather than a bath or changing the way you cook – can make a big difference to your safety if you have a seizure.

There is, however, another risk that you need to consider too. It’s very small, but significant, and it becomes more serious if you have frequent tonic-clonic seizures. It’s called Sudden Unexpected Death in Epilepsy – or SUDEP – and it’s something you may want to discuss with your medical team when you’ve read this leaflet.

What is SUDEP?

Each year, roughly one in every thousand people with epilepsy will die suddenly with no obvious cause. This is what we call SUDEP. The risk may be greater for a small number of people, which is why it’s important to talk to your doctors or nurses about your own condition. SUDEP usually occurs at night or during sleep. It’s often the case that the affected person will have had a seizure just before their death, although this isn’t universal.

Why does it happen?

Researchers are still investigating the causes of SUDEP and it’s unlikely there’s a single cause. During seizures, people may often see a change in their breathing or heart rhythm. Normally, this isn’t a problem, but if more serious changes occur – a major drop in oxygen levels in the blood, for example – then this might be a trigger for SUDEP.

Seizures may also affect the brain’s influence on breathing and the heart. Another possibility is that there is a genetic link between seizures and heart rhythm abnormalities. Advances in research mean that we’re getting closer to identifying the critical risk factors.

The strongest risk factor is a generalised tonic-clonic seizure (which some people call ‘grand mal’). The more frequent these convulsive seizures are, the higher the likelihood of SUDEP. As a result, the most important step you can take to avoid SUDEP is to minimise the number of seizures you have.

Read our Epilepsy. Be smart, be safe leaflet.

It gives a lot of practical advice about living with epilepsy that you can discuss with your family, friends and medical team.

Take your prescribed medication regularly and reliably.

Try a diary or smartphone app if you have trouble remembering.

Keep regular appointments with your doctor and attend your annual review.

Don’t make adjustments to your medication or stop taking it without talking to a doctor first. There is help available to safely look at your choices. If you’re making changes in your life – moving away from home or taking contraception for the first time – see if your epilepsy care plan needs to change.

Consider using EpSMon, a smartphone app to help you regularly assess and monitor your epilepsy, risks and wellbeing so you are aware of any changes. Find out more at www.epsmon.com. You could also tell your health professionals about the SUDEP and Seizure Safety Checklist, a communication tool to help them discuss and monitor your risks and wellbeing during your annual review. They can find out more at www.sudep.org/checklist.

How can you reduce your risk?

Identify the triggers for your seizures.

Do you find that your epilepsy is particularly affected by not sleeping enough, drinking too much alcohol or taking recreational drugs? If so, try to avoid the trigger.

Discuss other epilepsy treatments with your doctor.

If you feel your medication isn’t doing enough to control your seizures, there may be other options such as surgery.

Confide in your friends.

Tell them that you have epilepsy and explain what they should do if you have a seizure.

There is some evidence that it’s safer to be with someone when you have a seizure, so it might be worth discussing sharing a room with a family member if you are affected during your sleep. Alternatively, you might want to consider, with your doctor, a monitoring device that may alert them if you have a seizure.

‘Don’t make adjustments to your medication or stop taking it, without talking to a doctor first’