The Epilepsy Passport contains essential information about a child or young person’s epilepsy in an easy-to-carry format.

It aims to help children with epilepsy and their families communicate with healthcare professionals and to help healthcare professionals communicate with each other.

Ask your epilepsy specialist nurse or consultant about the Epilepsy Passport or visit:

www.epilepsypassport.org.uk
What is it?

The Epilepsy Passport contains essential information about a child’s epilepsy, including their medication history, emergency care plan and key professional contacts.

Why have it?

Studies have shown that communication between healthcare professionals and parents/carers of children with epilepsy can be difficult. The Epilepsy Passport has been designed to improve communication in outpatient settings and in an emergency.

Where is it available?

Epilepsy Passports are available at www.epilepsypassport.org.uk and must be filled in by a paediatrician or epilepsy nurse specialist at epilepsy clinic appointments. It will then be given to the child/young person and their parents/carers, who will carry it around and present it to healthcare professionals as and when it is needed.

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