Almost 1 in 5 (18%) women currently taking the epilepsy medication sodium valproate do not know it can potentially harm the development and physical health of their unborn child should they become pregnant. This was the worrying statistic revealed in a new survey carried out by three leading epilepsy charities.

More than 2,000 women with epilepsy took part in the survey, conducted by Epilepsy Action, Epilepsy Society and Young Epilepsy. The survey revealed that just over a quarter (28%) of women who responded, and are currently taking sodium valproate, had not been given information about risks for children exposed to the drug during pregnancy.

It is estimated that around 10% of babies born to women who take sodium valproate during pregnancy are born with physical disabilities. Up to 40% are at risk of developmental issues that can lead to learning difficulties.

These results are in spite of efforts by the Medicines and Healthcare products Regulatory Agency (MHRA) to raise awareness of the issue among healthcare professionals and women with epilepsy. In February 2016, the MHRA released a valproate toolkit to help healthcare professionals talk to women with epilepsy about the risks during pregnancy. The toolkit includes a credit card-sized patient card to be issued by pharmacists, booklets for healthcare professionals and women taking sodium valproate, and a checklist of important discussion points.

However, 18 months later, the survey shows that more than two thirds (68%) of respondents currently taking sodium valproate have still not received any of the toolkit materials.

Twenty-four hours after having her first baby at 16, Michelle Hackett started having multiple seizures and was prescribed sodium valproate. Less than a year later, her second baby was born with six toes on each of his feet, an underlying jaw and, when his teeth came through, eight of them were black and rotten.
Michelle says: “I had a scan and the hospital told me that because I was on Epilim (sodium valproate), I would have to get rid of the baby. They said it could be born with cerebral palsy. I refused. I asked why I hadn’t been told about the dangers around Epilim but they said because I was young, they didn’t think I would get pregnant again.

“I was very relieved Stephen didn't have cerebral palsy. I really feel he was one of the lucky ones but as a baby and toddler he had to have a lot of operations to correct his problems.

“I feel absolutely furious that I wasn't told about the side effects of Epilim. When I hear about mums with Epilim and what they go through it is heart-breaking. It is terrible to be given a drug for a severe illness and not be told about the side effects.

Michelle is now taking a different epilepsy medication and a year ago gave birth to a healthy baby girl, Thea.

Epilepsy Action, Epilepsy Society and Young Epilepsy are now calling on the government to ensure that repeat prescriptions for sodium valproate for women and girls of childbearing age are not routinely renewed for more than 12 months without a face-to-face consultation with a doctor or nurse. This consultation must include personal and tailored information about the risks around sodium valproate during pregnancy. Information should also be provided in written format.

This will help to ensure that women and girls with epilepsy who are prescribed sodium valproate are fully informed about the risks to their unborn baby should they continue to take the drug while pregnant.

Clare Pelham, chief executive at Epilepsy Society said: ‘This is a very simple and straightforward ask of the government. No additional funding is required for the NHS. All we are asking is that doctors and nurses have a yearly conversation with women and girls who are taking sodium valproate, to make absolutely certain they are aware of the risks associated with the drug during pregnancy.

‘We know that for some women, sodium valproate will be the only drug that controls their seizures and they must carefully weigh the benefits against the risks. But what is unforgiveable is for women not to be informed of those risks. They must be able to make an informed choice about their treatment in conjunction with their doctor.’

Simon Wigglesworth, deputy chief executive at Epilepsy Action, said: “It is vital that women with epilepsy get the right information about their care and treatment to ensure a healthy pregnancy and minimise the risks associated with sodium valproate.

“Yet these figures suggest that information is not filtering down to women and those conversations about the potential risks are not always happening. Discussions with a health professional about these risks should be a mandatory part of care for all women with epilepsy so they can make informed choices, ideally before they conceive.”

Carol Long, Chief Executive of Young Epilepsy, said: “Epilepsy continues to be a ‘Cinderella-like’ condition and it is important that women who may be planning to have a family receive the right advice at the right time; to prevent their unborn babies being put at risk.
“Sodium valproate can provide life-changing support for many young people with epilepsy. It is the third most-prescribed anti-epilepsy medication, however, the percentage of women who do not know the risks of taking the drug during pregnancy is far too high. That information must be made clearer, and medical professionals must be given more support to understand and be proactive in flagging the risks. Women must feel more empowered to discuss the issue with their epilepsy specialist, so that they can make a more informed decision about their future.”

The survey is published in advance of the European Medicines Agency's (EMA) public hearing into sodium valproate next Tuesday, 26 September. The hearing, in London, will be the first time the EMA has held a public hearing as part of the safety review of a drug.

The three charities will be presenting the survey results at the public hearing along with patient groups, healthcare professionals and other interested bodies from across Europe.

For advice and information about epilepsy and pregnancy, visit the charities’ websites: epilepsy.org.uk, epilepsysociety.org.uk and youngepilepsy.org.uk

- ENDS -

Notes to editor:

1. Epilepsy Action, Epilepsy Society and Young Epilepsy surveyed 2,000 women between the ages of 16 and 50 in the UK. The survey is currently running. This data was collected between 17 August and 20 September 2017. The survey has been held in conjunction with the MHRA.

2. Around one in 100 people in the UK has epilepsy. Epilepsy is the tendency to have recurrent seizures. There are around 60 different types of epileptic seizure, and everyone experiences them differently. They range from experiencing strange sensations to falling unconscious, but all have a major impact on daily life. With the right treatment, 70% of people could be seizure-free.

3. For media enquiries:
   Epilepsy Action: Louise Cousins – press@epilepsy.org.uk or 0113 210 8870
   Epilepsy Society: Nicola Swanborough - Nicola.Swanborough@epilepsysociety.org.uk or 01494 601 417
   Young Epilepsy: John Price - JPrice@youngepilepsy.org.uk or 01342 832243 ext 505