Commentary
GlaxoSmithKline (GSK) put forth their final results of the International Lamotrigine Pregnancy Registry, reporting major congenital malformations (MCMs) in 35 infants out of 1,558 first trimester monotherapy exposures, a frequency of 2.2%.
(95% confidence interval [CI] of 1.6–3.1%) (1). This international registry has been enrolling since 1992, with 65% of subjects enrolled from the United States (Lamictal was approved in the United States in late 1994), and 20% more from northern Europe. The enrollment was closed in June 2009, since the proscribed enrollment goal of 1,000 first-trimester lamotrigine exposures was met. This enrollment target was thought to provide adequate power to identify a large increase in the frequency over the expected rate of all combined MCMs with lamotrigine exposure.

In the early 1990s, Glaxo Wellcome launched five international pregnancy registries: 1) the Acyclovir Pregnancy Registry, 2) the Antiretroviral Pregnancy Registry, 3) the Lamotrigine Pregnancy Registry, 4) the Sumatriptan Pregnancy Registry, and 5) the Bupropion Pregnancy Registry, with the aim of obtaining information about rates of teratogenicity of these compounds (2). The International Lamotrigine Pregnancy Registry has been lauded by the epilepsy community (although it enrolled subjects taking lamotrigine for reasons other than epilepsy); it was started early in the marketed use of lamotrigine, and therefore was definitely a departure from the postmarketing activities for other newer antiepileptic drugs (AEDs) for which teratogenic risk in humans was an afterthought or completely ignored. In 2002, with 9 years of data, GSK published their preliminary results (3). (An even more preliminary report was published in 2000 of the results of all five registries above (4).) They reported the occurrence of three infants with MCMs (1.8%) out of 168 first-trimester lamotrigine exposure (95% CI 0.5–5.5%) (3).

In 2005, in another update, GSK reported that of 414 first-trimester exposures to lamotrigine monotherapy, 12 outcomes with MCMs occurred (2.9%, 95% CI 1.6–5.1%) (5). Another analysis was published in 2007, in which the frequency of MCMs in 802 first-trimester monotherapy lamotrigine exposures was 2.7% (95% CI 1.8–4.2%) (6). The main point of this particular manuscript, however, was to document the absence of a dose–dependent relationship with MCMs, which was analyzed by logistic regression using 100-mg increments, up to 400 mg per day. This GSK publication appeared to be in response to a report from the U.K. Pregnancy Register in 2006 (7), in which a lamotrigine dose–dependent relationship with MCMs between the use of less versus more than 200 mg per day was found. An increased risk of isolated facial clefts in exposed infants was pointedly not found in the final GSK registry report, refuting what was first reported by the North American AED Pregnancy Registry (8) but later not borne out in a case control study from the EUROCAT database (9).

The frequency of malformations in the GSK registry has basically been low at all assessment points during the registry, from 1.8 to 2.9 percent, with the final number at 2.2% surrounded by very narrow confidence intervals, which have also gradually been closing in around this number over the years, now at 1.6 to 3.1 percent. However, this confidence interval means that it is just as likely that the “true” frequency of malformations can be 3.1% or 1.6% as it is 2.2%. The authors compare this result to MCM frequencies in the general population from large surveillance databases, of 1.6 to 2.2 percent. The GSK results are similar to findings from the EURAP registry, in which the frequency of MCMs with first-trimester lamotrigine monotherapy exposure was 2.3%, with 30 infants of 1,280 exposures having MCMs at 2 months follow-up; seven additional MCMs were found in this cohort after 1 year of follow-up (10). Therefore, the frequency of malformations in the GSK registry is consistent over 18 years, very close to the general population and nearly identical to that of a concurrently reported academic database. The internal consistency and favorable comparisons to external databases support the validity of the GSK registry findings.

Of interest, another very recent report from a Danish population-based database found a frequency of MCMs of 3.7% in infants exposed to lamotrigine in the first trimester (38 of 1,019 exposures) (11). After the authors adjusted for the diagnosis of epilepsy, the risk of MCMs for all AED exposures was decreased; overall, the presence of an epilepsy diagnosis itself contributed about a 40 to 50 percent increased frequency of birth defects compared with the non-AED-exposed comparator group. Further, after adjusting for an epilepsy diagnosis, the risk of MCMs with lamotrigine exposure compared with controls was not significant. This may be the best data thus far to quantitate a risk imparted by epilepsy itself.

Limitations of the GSK study methodology include the risk of reporting bias, since women were referred to the registry by a health care provider and enrolled voluntarily, whether they knew any ultrasound results or not at the time of enrollment. It is unclear if this enrollment methodology would bias the study toward a lower MCMs frequency. The results would suggest not; there were three cases of anencephaly reported, and this malformation is reliably seen on ultrasound between 10 and 14 weeks of pregnancy. This surprising finding and its association with lamotrigine should be further assessed using the EUROCAT data as well. The GSK registry enrollment strategy is similar to that of the EURAP registry; neither registry is population representative and may even include some common subjects. The low frequency of MCMs in the GSK final report is consistent with the stated methodology of including birth defects found only before hospital discharge. This frequency of malformations is identical to that found in the North American Pregnancy Registry strategy, which also uses a very short period after birth to assess MCMs, but is shorter than the EURAP assessments, which are at 2 months and up to 1 year. Although there was no concurrent control group in the GSK study, the data within the cohort are revealing and consistent; a higher frequency of birth defects with polytherapy that includes valproate has been consistently reported, now at 10.7% in this final report.

The issue of a lamotrigine dose–dependent association with MCM frequency is not sorted out. The GSK registry did not detect it; however, the EURAP report showed an increase in MCMs with doses ≥300 mg per day. It is intuitive that a greater exposure would pose a greater risk, and this is certainly present for valproate exposure. This relationship is more clearly detected with valproate because the frequency of adverse events is in general much greater.

GSK has positioned lamotrigine as an AED to consider when treating women of childbearing potential. In spite of the challenges around the variable clearance of lamotrigine with exogenous hormones such as oral contraceptives and with endogenous hormones during pregnancy, the GSK pregnancy
registry and other registries bear out a low frequency of MCMs with use during pregnancy. This is good and useful news for patients, healthcare providers, and for GSK. Did we need an industry-sponsored pregnancy registry to find this out? These findings would have eventually been reported in other registries. However, putting aside the marketing strategy, this registry was established before any of the academic registries, which, outside of the national health registries are supported by industry funding in any case. Even the Danish database study cited herein was supported in part by Lundbeck. The GSK registry was established to answer a dire clinical question, provided early reports on frequencies of malformations, was not more biased than other registries, and of course helped to market Lamictal. Best answer D.

by Cynthia L. Harden, MD

References

American Epilepsy Society
Epilepsy Currents Journal
Disclosure of Potential Conflicts of Interest

Instructions
The purpose of this form is to provide readers of your manuscript with information about your other interests that could influence how they receive and understand your work. Each author should submit a separate form and is responsible for the accuracy and completeness of the submitted information. The form is in four parts.

1. Identifying information.
   Enter your full name. If you are NOT the main contributing author, please check the box “no” and enter the name of the main contributing author in the space that appears. Provide the requested manuscript information.

2. The work under consideration for publication.
   This section asks for information about the work that you have submitted for publication. The time frame for this reporting is that of the work itself, from the initial conception and planning to the present. The requested information is about resources that you received, either directly or indirectly (via your institution), to enable you to complete the work. Checking “No” means that you did the work without receiving any financial support from any third party – that is, the work was supported by funds from the same institution that pays your salary and that institution did not receive third-party funds with which to pay you. If you or your institution received funds from a third party to support the work, such as a government granting agency, charitable foundation or commercial sponsor, check “Yes”. Then complete the appropriate boxes to indicate the type of support and whether the payment went to you, or to your institution, or both.

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2. First Name Cynthia     Last Name Harden  Degree MD

3. Are you the Main Assigned Author?  ☑ Yes  ☐ No 

   If no, enter your name as co-author:

4. Manuscript/Article Title: Final results from the lamotrigine international pregnancy registry

5. Journal Issue you are submitting for:  Epilepsy Currents

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