Associations Between ADHD Medication Use in Pregnancy and Severe Malformations Based on Prenatal and Postnatal Diagnoses: A Danish Registry-Based Study

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ABSTRACT

Objective: Attention-deficit/hyperactivity disorder (ADHD) medications are increasingly used in pregnancy. Studies on the pregnancy safety of these medications that are restricted to live births may underestimate severe teratogenic effects that cause fetal demise or termination of pregnancy. The present study addresses this limitation by including data from both prenatal and postnatal diagnoses of major malformations.

Methods: A nationwide registry-based study was conducted of 364,012 singleton pregnancies in Denmark from November 1, 2007, to February 1, 2014. Exposures to ADHD medication were obtained from redeemed prescriptions from the Danish Health Services Prescription Database. Outcome data included prenatally diagnosed malformations from the Danish Fetal Medicine Database and postnatally diagnosed malformations from the Danish National Patient Registry. The primary outcome was major malformations overall, and secondary outcomes were malformations of the central nervous system and cardiac malformations. The comparison group was pregnancies with no redeemed prescriptions for ADHD medication. We defined severe cardiac malformations (SCM) as concurrent diagnoses of a cardiac malformation with miscarriage, termination, stillbirth, postnatal death, or cardiac surgery within 1 year of birth.

Results: The prevalence of first-trimester exposure to ADHD medication increased during the study period from 0.05% in 2008 to 0.27% in 2013, with the majority (473/569) of the exposures being to methylphenidate. There were 5.1% malformations overall and 2.1% cardiac malformations among the exposed compared to 4.6% and 1.0%, respectively, among the unexposed. For methylphenidate, the adjusted prevalence ratios (PRs) were 1.04 (95% confidence interval [CI], 0.70–1.55) for malformations overall and 1.65 (95% CI, 0.89–3.05) for any cardiac malformations (number needed to harm [NNH] = 92), with septum defects in 10 out of 12 cases. The PR for ventricular septal defect was 2.74 (95% CI, 1.03–7.28) and for SCM, 2.59 (95% CI, 0.98–6.90).

Conclusions: Exposure to methylphenidate was not associated with an increased risk of malformations overall in data that included information from both prenatal and postnatal diagnoses of major malformations. There was an increased risk of cardiac malformations with NNH of 92 based on 12 cases among the exposed. More data are needed on other types of ADHD medication.

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Clinical Points

- Use of ADHD medication increased 5-fold in the study period, underscoring the need for updated and unbiased information on pregnancy safety.
- There was no association between methylphenidate and malformations overall in a study with information on both prenatal and postnatal diagnoses.
- An increased risk of cardiac malformations (number needed to harm = 92) based on 12 cases among the exposed was found.

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Use of attention-deficit/hyperactivity disorder (ADHD) medication in pregnancy increased from 5 per 100,000 person-years in 2003 to 533 per 100,000 in 2010 in Denmark. Such dramatic increases have been seen in other populations even though there is a relative paucity of pregnancy safety data for ADHD medication. In fact, most ADHD medications are sympathomimetics with pharmacologic actions similar to those of known teratogens (amphetamine and cocaine), which further underscores the need for safety data on ADHD medication exposure in pregnancy.

Reassuringly, a 2018 meta-analysis of 8 cohort studies did not find an association of exposure to any ADHD medication during pregnancy and any major malformation (risk ratio [RR] = 1.05; 95% confidence interval [CI], 0.96–1.16) or cardiac malformations (RR = 1.08; 95% CI, 0.92–1.27). However, methylphenidate was associated with an increased risk of cardiac malformations (RR = 1.27; 95% CI, 0.99–1.63), with the result nearly solely based on data from a large study from the United States and the Nordic countries that included 3,474 methylphenidate-exposed pregnancies.

The above evidence, similarly to most such studies, stems from data on pregnancies ending in live births. Such studies cannot detect associations for severe malformations that cause spontaneous or induced pregnancy loss. Lack of data on malformations ending in a pregnancy loss may consequently lead to an underestimation of the true occurrence of malformations and may result in survivor bias if only data on liveborn children are available.

We assembled pregnancies with complete information on outcomes, ie, data on malformations detected by prenatal ultrasound, terminations, miscarriages, stillbirths, and malformations among liveborn children, and examined the potential association between first-trimester exposure to ADHD medication and major malformations, overall and specifically malformations of the central nervous system and heart.

METHODS

This prevalence study was set in Denmark, a prosperous country with universal access to health care. We used routinely collected data from 5 nationwide databases: the Danish Fetal Medicine Database, the Danish National Patient Registry, the Danish Medical Birth Registry, and the Danish Health Services Prescription Database, all linked on individual level (including mother-child linkage) via a unique identifier from the Danish Civil Registration System.

The study population comprised all clinically recognized singleton pregnancies with a live fetus at the first-trimester scan from 11 gestational weeks, with conception dates from November 1, 2007, through February 1, 2014. Pregnancies terminated before gestational week 22 were identified in the Danish National Patient Registry, and pregnancies ending in live birth or stillbirth from week 22 onward were identified from the Danish Medical Birth Registry. Gestational age at pregnancy end was estimated from ultrasound measurements in early pregnancy or, if unavailable, calculated from the last menstrual period. Conception date was estimated by subtracting the gestational age from the date of pregnancy end and adding 14 days. Pregnancies with fetal chromosomal abnormalities were excluded regardless of presence of other malformations.

Exposure to ADHD medication was measured using redeemed prescriptions through linkage to the Danish Health Services Prescription Database. First-trimester exposure was defined as 1 or more redeemed prescriptions from 28 days before conception date through 70 days after conception date. We restricted the cohort by excluding preconception exposure to ADHD medication (former use), an adapted model previously suggested by Huybrechts et al and Jimenez-Solem et al. The unexposed group had no recombinations for an ADHD medication from 28 days before conception date through 70 days after conception date (first trimester) and from 365 to 183 days before conception date (former use). Medications were analyzed as any ADHD medication and as specific drugs (methylphenidate, modafinil, atomoxetine).

Major malformations were classified according to the European Surveillance of Congenital Anomalies (EUROCAT) classification, version 2014. Data on prenatally diagnosed major malformations came from the Danish Fetal Medicine Database, which contains data on malformations diagnosed prenatally including all first-trimester screenings and second-trimester malformation...
scans starting from gestational week 11. The database was 95% complete during its period of coverage. The primary outcome was major malformations overall, and secondary outcomes were malformations of the central nervous system or the heart. Cardiac malformations were classified by severity into severe cardiac malformations (SCM; malformations in pregnancies ending in a miscarriage or termination, stillbirth, death, or cardiac surgery during the first year of life, arranged in hierarchy of termination > miscarriage > death > surgery) and non-SCM (all other cardiac malformations). Indication for terminations after week 12 was either on fetal indication (eg, malformation) or non-fetal indication (eg, severe disease of the pregnant woman).

Data on covariates, obtained from the above-mentioned data sources, included maternal ethnicity (Caucasian, other), civil status (living with partner, yes/no), parity (nulliparous/multiparous), age at conception (≤ 35, > 35 years), prepregnant body mass index (BMI; < 25, 25 to < 30, ≥ 30 kg/m²), smoking during pregnancy, redeemed prescription of a known teratogen from 180 days preconception and through the first trimester, first-trimester and 2 years preconception dispensing of other medication (antihypertensives, antidiabetic agents, antiepileptics, anxiolytics, hypnotics, and sedatives), and maternal hospital diagnoses of ADHD or diabetes 2 years pre–conception date.

We tabulated maternal characteristics according to first-trimester exposure to ADHD medication. We used log-binomial regression to estimate prevalence ratios (PRs) with 95% CIs for the study outcomes. We presented crude PRs; PRs minimally adjusted, by including in a regression model, for age, smoking, and ADHD diagnosis (if n > 5 cases); and PRs fully adjusted using propensity-score fine stratification (50 strata) for ethnicity, civil status, parity, age, BMI, smoking, exposure to teratogens, antihypertensives, antidiabetics, use of other psychotropic drugs, ADHD diagnosis, and diabetes diagnosis.

In sensitivity analyses, we redefined first-trimester exposure by 2 or more redeemed prescriptions to assess the effect of misclassification on the results. We repeated the analyses restricting the study population to pregnancies ending in live births to assess the impact of the resulting selection bias on the results.

Statistical analyses were performed with Stata (version 14, StataCorp, Texas) and SAS (version 9.4, SAS Inc, Cary, North Carolina). In all analyses, cell counts under 5 were masked to avoid identification of individuals.

RESULTS

We identified 366,489 eligible pregnancies during the study period. After excluding 354 pregnancies due to coding errors and 2,123 pregnancies with chromosome anomalies, the study population comprised 364,012 clinically recognized singleton pregnancies in week 11 onward, of which, from week 12 and onward, 96% had data from ultrasound examinations from the Danish Fetal Medicine Database.

The prevalence of first-trimester exposure to ADHD medication was 0.16% (n = 569) on average. It increased more than 5-fold during the study period, from 0.05% in 2008 to 0.27% in 2013 (Figure 1). The majority of the exposed pregnancies (473/569, 83%) were exposed to methylphenidate. Only half of the women exposed to ADHD medication had a hospital diagnosis of ADHD 2 years prior to the conception date. Some of the remaining exposed pregnancies may be because the primary hospital diagnosis was set earlier than 2 years prior to conception, and the treatment was maintained by the general practitioner. Compared with unexposed women, women who used ADHD medication were younger and more likely to smoke and/or use other psychotropic medications (Table 1). Termination for any reason after gestational week 11 had a prevalence of 9.8% among pregnancies with first-trimester exposure to ADHD medicine and 2.6% among the unexposed (Table 2). The majority of terminations (after week 12) in the exposed group were due to nonfetal indications.

The overall prevalence of malformations was 5.1% among the methylphenidate-exposed pregnancies and 4.6% among the unexposed pregnancies. The prevalence of cardiac malformations was 2.1% among the methylphenidate exposed and 1.0% among the unexposed. Number needed to harm (NNH) was 92. For SCM, the prevalences were 0.9% among the methylphenidate exposed and 0.2% among the unexposed.

Most PR estimates became attenuated by improved control of confounding (Table 3). For methylphenidate, the fully adjusted PRs were 1.04 (95% CI, 0.70–1.55) for malformations overall, 1.65 (95% CI, 0.89–3.05) for cardiac malformations, and 2.59 (95% CI, 0.98–6.90) for SCM. In the pregnancies with cardiac malformations, 10 out of 12...
Table 1. Characteristics of Women Exposed to ADHD Medication During First Trimester and Unexposed

<table>
<thead>
<tr>
<th>Age at conception, mean (SD) y</th>
<th>Exposed (n = 569)</th>
<th>Unexposed (n = 363,219)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parity</td>
<td>26.4 (5.7)</td>
<td>29.7 (5.1)</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 25</td>
<td>259 (45.5)</td>
<td>212,479 (58.5)</td>
</tr>
<tr>
<td>25–29.9</td>
<td>106 (18.6)</td>
<td>68,264 (18.8)</td>
</tr>
<tr>
<td>≥ 30</td>
<td>128 (22.5)</td>
<td>60,934 (16.8)</td>
</tr>
<tr>
<td>Smoking in pregnancy</td>
<td>208 (36.5)</td>
<td>36,699 (10.1)</td>
</tr>
<tr>
<td>Civil status, living with partner</td>
<td>498 (87.5)</td>
<td>327,267 (90.1)</td>
</tr>
</tbody>
</table>

Table 2. Pregnancy Outcome for Exposure to ADHD Medication During First Trimester and Unexposed

<table>
<thead>
<tr>
<th>Termination ≥ week 11 + 0</th>
<th>Exposed (n = 569), n (%)</th>
<th>Unexposed (n = 363,219), n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Miscarriage &lt; week 22 + 0</td>
<td>56 (9.8)</td>
<td>9,333 (2.6)</td>
</tr>
<tr>
<td>Stillborn, singleton ≥ week 22 + 0</td>
<td>&lt; 5a</td>
<td>1,124 (0.3)</td>
</tr>
<tr>
<td>Live born</td>
<td>492 (86.5)</td>
<td>341,688 (94.1)</td>
</tr>
</tbody>
</table>

Major malformations

- Any: 29 (5.1) vs. 16,857 (4.6)
- Cardiac: 12 (2.1) vs. 3,734 (1.0)
- Central nervous system: < 5a, 1,259 (0.4)
- Eye: 0 vs. 494 (0.1)
- Respiratory: < 5a, 673 (0.2)
- Orofacial clefts: 0 vs. 629 (0.2)
- Face: 0 vs. 341 (0.1)
- Digestive system: < 5a, 1,088 (0.3)
- Abdominal wall defect: < 5a, 212 (0.1)
- Urinary: < 5a, 2,659 (0.7)
- Genital: < 5a, 1,274 (0.4)
- Limb: 5 (0.9) vs. 4,689 (1.3)

*Number is rounded to nearest 10 or reported as ≤ 5 to comply with data protection regulations.

Abbreviation: ADHD = attention-deficit/hyperactivity disorder.

We defined SCM as termination, miscarriage, death, or surgery. Previous studies did not consider severity; however, we find this outcome to be of major importance. Clinically, it is important because both health care providers and patients ask about the severity of outcomes. Methodologically, it is important because the classification can generate power by assembling rare outcomes and reduce the risk of surveillance bias in the severe group through the robust outcomes of death and surgery. This might contribute to the stronger association between methylphenidate and SCM than between methylphenidate and any cardiac malformations.

The results are in accordance with earlier studies, including a recent systematic review that reported a 1.3-fold association between first-trimester methylphenidate exposure and prevalence of overall cardiac malformation.4 However, our outcomes, according to the EUROCAT definition of major cardiac malformations, did not include patent ductus arteriosus and patent foramen ovale. The large study of Huybrechts et al4 included in the primary analysis women from the United States and replicated analyses adding pregnancy registry data from 5 Nordic countries (Denmark, Finland, Iceland, Norway, and Sweden). In this pooled analysis, the study found a 28% increase in cardiac malformations after methylphenidate exposure; furthermore, after patent ductus arteriosus and patent foramen ovale were excluded, the associations strengthened for methylphenidate: OR = 1.50 (95% CI, 1.05 to 2.14), in accordance with our findings.4 Moreover, similarly to earlier studies, we did not find an association between ADHD medication and overall malformations.1,4,18–20 Similarity of results is expected given the overlap of our study population with the studies by Harvig et al1 (overlap birth years 2008, 2009) and Pottegård et al20 (overlap birth years 2008–2012) and the Danish population in the study by Huybrechts et al4 (overlap birth years 2008–2011), even with our use of prenatal data, as only 1 exposed case was lost prenatally. In this study, inclusion of

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exposed cases were septum defects, atrial septal defect (ASD) or ventricular septal defect (VSD), yielding fully adjusted PRs in methylphenidate-exposed pregnancies of 1.21 (95% CI, 0.84–1.65) for ASD and 2.74 (95% CI, 1.03–7.28) for VSD. The fully adjusted PR for CNS malformations was 2.03 (95% CI, 0.65 to 6.27). The estimates for SCM and CNS malformations were based on few exposed cases.

Sensitivity analyses, with exposure defined by 2 or more redeemed prescriptions during the first trimester, resulted in a decrease of cases of cardiac malformations among the exposed from 12 to 9. For methylphenidate, the fully adjusted PR for cardiac malformation was 1.75 (95% CI, 0.84–3.65) (data not shown). In analyses restricted to pregnancies ending in live births, the results were unchanged for any malformations and cardiac malformations, but crude estimates strengthened in cardiac malformation subgroups and CNS malformations, all based on few exposed cases (data not shown).

DISCUSSION

In this population-based study, methylphenidate was associated with a 1.65-fold increased risk of cardiac malformations, 2.74-fold increased risk of VSD, and 2.59-fold increased risk of SCM. Septal defects were present in 10 of 12 cases of exposed cardiac malformations.
prenatal data added 233 cases of cardiac malformations that were not in live births (terminated, miscarriage, or stillborn). Only 1 case was exposed to ADHD medication, resulting in the estimates of cardiac malformations being unchanged.

Similar to previous findings,\textsuperscript{1,19} we found that pregnancies exposed to ADHD medication were 4 times more likely to end in a termination of pregnancy than unexposed pregnancies. In addition, a Danish study highlighted the potential for confounding by indication, as the authors found that exposure to ADHD but not to ADHD medications was associated with both spontaneous abortion and preterm birth.\textsuperscript{19}

The estimates of exposure prevalence were similar to those in previous studies in the first conception years (eg, 0.05% in 2008),\textsuperscript{1,4} and the 5-fold increase is in continuation of the previously described increased use of ADHD medication.\textsuperscript{1} The prevalences of malformations per 1,000 singleton pregnancies (46 vs previously 35 per 1,000)\textsuperscript{4} and of cardiac malformations (10 vs previously 13 per 1,000),\textsuperscript{4} after excluding chromosomal anomalies, were in accordance with previous findings. The degree of selection bias was minimal in our study owing to the completeness and nationwide coverage of the data sources, allowing inclusion of nearly all pregnancies and all diagnosed malformations both prenatally and postpartum within 1 year.

A limitation of the study is the possibility of confounding by indication, even though several approaches were used to adjust for or assess the extent of confounding. Atomoxetine, which in contrast to methylphenidate is a nonstimulant ADHD drug, may have been an ideal negative control which in contrast to methylphenidate is a nonstimulant ADHD drug, may have been an ideal negative control.
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REFERENCES
1. During this study period from 2008 to 2013, how did the prevalence of ADHD medication use during pregnancy change?
   a. Unchanged
   b. Moderately decreased
   c. Dramatically decreased
   d. Increased

2. Lottie is a 28-year-old woman who is planning her first pregnancy. She was diagnosed with ADHD and has been treated with methylphenidate for the last 7 years. Lottie asks you if the medication is going to be harmful for the unborn child. Which statement reflects your evidence-based answer?
   a. Methylphenidate is not harmful for the unborn child.
   b. Methylphenidate is associated with an increased risk of any major malformation.
   c. Methylphenidate is associated with an increased risk of cardiac malformations.
   d. Methylphenidate is associated with an increased risk of severe central nervous system malformations.

3. What is the challenge related to evaluating drug safety in only live births?
   a. To underestimate an association between drug exposure and teratogenic effects
   b. To overestimate the risk of teratogenic effects with drug exposure
   c. Nothing; there is no risk of changing the estimate of teratogenic effects because so few births are not live.
   d. To create a false association between drug exposure and teratogenic effects