Cardiovascular safety of methylphenidate among children and young people with attention-deficit/hyperactivity disorder (ADHD): nationwide self controlled case series study

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ABSTRACT

OBJECTIVE
To determine whether treatment with methylphenidate in children and young people with attention-deficit/hyperactivity disorder (ADHD) was associated with cardiovascular events.

DESIGN
Self controlled case series analysis.

SETTING
Nationwide health insurance database, 1 January 2008 to 31 December 2011, in South Korea.

PARTICIPANTS
1224 patients aged ≤17 who had experienced an incident cardiovascular event and had had at least one incident prescription for methylphenidate.

MAIN OUTCOME MEASURES
A recorded diagnosis (either a primary or secondary cause) of any of the following cardiovascular adverse events: arrhythmias (ICD-10 (international classification of diseases, 10th revision) codes I44, I45, I47-I49), myocardial infarction (code I21), ischemic stroke (code I63), or heart failure (code I50). Incidence rate ratios were calculated with conditional Poisson regression and adjusted for time varying comorbidity and comedication.

RESULTS
Increased risk of arrhythmia was observed in all exposed time periods—that is, periods of treatment with methylphenidate—(incidence rate ratio 1.61, 95% confidence interval 1.48 to 1.74), and the risk was highest in the children who had congenital heart disease. No significant risk of myocardial infarction was observed for all exposed time periods (1.33, 0.90 to 1.98), though risk was higher in the early risk periods between eight and 56 days after the start of treatment with methylphenidate. No significant increased risk was observed for hypertension, ischemic stroke, or heart failure.

CONCLUSION
The relative risk of myocardial infarction and arrhythmias is increased in the early period after the start of methylphenidate treatment for ADHD in children and young people. Though the absolute risk is likely to be low, the risk-benefit balance of methylphenidate should be carefully considered, particularly in children with mild ADHD.

Introduction

Drugs to treat attention-deficit/hyperactivity disorder (ADHD) have been shown to be efficacious in reducing symptoms of impulsivity and hyperactivity in children, but concerns have been expressed about possible adverse cardiovascular events with the first line treatment, methylphenidate. In 2009, one case-control study reported a strong association between use of stimulants in young people and sudden death (1.8%, 10 users among 564 cases) compared with use of stimulants in death in people in motor vehicle crashes (0.4%, 11 users among 564 cases) (odds ratio 7.4, 95% confidence interval 1.4 to 74.9). Comparability between the cases and controls, infrequent exposure, and potential recall bias were limitations in this analysis.

Subsequently, large retrospective population based cohort studies have found no evidence of an increased risk of myocardial infarction or stroke associated with methylphenidate. A systematic review reported that findings of observational studies of children and adolescents were variable and suggested that statistical power was a limitation as the absolute risk of cardiovascular events was low in the populations studied and the study designs meant that confounding could not be ruled out. Given this uncertainty around the risks associated with methylphenidate, case reports of adverse cardiovascular events continue to be published. There have been five case reports of life threatening heart failure associated with methylphenidate reported from Sweden. Additionally, small but significant increases in blood pressure have been reported in children and adolescents. Despite these latter reports, no observational studies have been performed to examine the risk of heart failure or hypertensive disease in children and adolescents exposed to methylphenidate.

We explored whether methylphenidate use is a triggering factor among children who experience cardiovascular events—including arrhythmia, hypertension, myocardial infarction, ischemic stroke, or heart failure—using a self controlled case series study design. Using this technique, which is a within person design, we looked at the incidence of each cardiovascular adverse event in children during periods in which they...
were exposed to methylphenidate compared with periods in which they were not exposed.

**Methods**

**Database**

We used the South Korea national health insurance claims database. The national health insurance program was initiated in Korea in 1977 and achieved universal coverage of the entire population by 1989. The database contains all information on the diagnoses and prescribed drugs for about 50 million Koreans. We obtained the claims data for children and young people aged ≤17 with diagnosis of ADHD (ICD-10 (international classification of diseases, 10th revision) code F90) that had been submitted by healthcare providers from 1 January 2007 to 31 December 2011. The claims database included an anonymized identifier representing each individual together with age, sex, diagnoses, and prescription drugs. Information on prescribed drugs included generic name, prescription date, duration, and route of administration.

All diagnoses are coded according to ICD-10. Previous validation studies have compared the diagnoses derived from the claims database with the ideal of actual diagnoses recorded in the patients’ medical records obtained from hospital or clinic chart review. The studies reported that the positive predictive value of the diagnoses was 83.4% for ischemic stroke (code I63) and 70% for myocardial infarction (code I21). The overall positive predictive value of all diagnoses was about 70%.

**Self controlled case series design**

We used a self controlled case series design in this analysis. This design is derived from the cohort design and determined the incidence of cardiovascular adverse events during periods in which the participants were exposed to methylphenidate compared with the incidence in periods in which they were not exposed. We used a bi-directional study design and collected data on cardiovascular adverse events in unexposed time either before and after exposure. The study design minimizes confounding because the patients each act as their own control. This method therefore controls for known and unknown confounders specific to the individual patient that do not vary over time, such as sex and genetic factors. The method enables comparisons in a population of individuals who have both the adverse event and exposure of interest. This design can be particularly useful when secondary healthcare data are used for pharmacoepidemiological research when limited information is available on potential confounders.

![Fig 1](https://www.bmj.com/)

**End of observation**

31 Dec 2011

**Start of observation**

1 Jan 2008

**Bi-directional case series study**

**Unexposed**

**Exposed to methylphenidate**

**Event**

**Fig 1** Design of self controlled case series study in children and young people with ADHD and cardiovascular adverse events

**Patient involvement**

No patients were involved in setting the research question or the outcome measures, nor were they involved in developing plans for design or implementation of the study. No patients were asked to advise on interpretation or writing up of results. There are no plans to disseminate the results of the research to study participants or the relevant patient community.

**Participants**

All data used were obtained from secondary electronic records for the study participants. These were all patients aged ≤17 who had at least one recorded diagnosis of ADHD (ICD-10 code F90), had started taking methylphenidate (ATC (Anatomical Therapeutic Chemical) code N06BA04), and had an incident cardiovascular adverse event with a recorded diagnosis during the study period (1 January 2008 and 31 December 2011). Patients were considered as new users of methylphenidate if they had not received a prescription during the preceding year.

Cardiovascular adverse events were defined as a recorded diagnosis (primary or secondary) for arrhythmias (codes I44, I45, I47, I48, I49), hypertension (code I10-I15), myocardial infarction (code I21), ischemic stroke (code I63), or heart failure (code I50). For patients with multiple diagnoses recorded, we included only the first event to avoid potential bias of the second event being influenced by the first event. The observation period started on 1 January 2008 and ended on 31 December 2011. There were five cases of death after the diagnosis of a cardiovascular adverse event, and for these cases we censored follow-up time at date of death.

**Assessment of exposure**

The index date was defined as the first date of prescription of methylphenidate. We calculated the length of exposure using information on prescription date and prescribed days in the database. Person time was divided into methylphenidate exposure risk periods: 13 days, 4-7 days, 8-14 days, 15-28 days, 29-56 days, and >56 days after prescription of the drug. Patients contributed to consecutive exposure risk periods while they were continuously exposed to the drug. All remaining time before and after exposure was considered as unexposed.
We included the following time varying covariates in the model: age, comorbidities (depressive episode (codes F32-F33, F34.1, F41.2), tic disorders (code F95), emotional disorders with onset specific to childhood (code F93), conduct disorders (code F91), manic episodes (code F30), bipolar affective disorders (code F31), and mental retardation (codes F70-79) and use of comedications (atomoxetine, antipsychotics (ATC code N05A), antidepressants (code N06A), anxiolotics (code N05B), antiepileptics (code N03A), and anticholinergic agents (code N04A)). Atomoxetine, a non-stimulant treatment for ADHD, is approved for secondary treatment. We could not find any participants taking just atomoxetine—that is, all records of atomoxetine were prescriptions with methylphenidate.

**Statistical analysis**

We determined the duration of exposure in the study period and calculated incidence rates for cardiovascular events in each of the exposure risk periods. Incident rate ratios in periods of exposure to methylphenidate compared with unexposed periods were calculated with conditional Poisson regression. We estimated the adjusted incidence rate ratios and their 95% confidence intervals for exposure overall and for each predetermined exposure risk period. Incidence rate ratios were calculated for each outcome event separately.

Additional analysis was performed according to the presence of congenital heart disease (ICD-10 Q20-Q24), and we also conducted a subgroup analysis according to the daily prescribed dose of methylphenidate (low and high). The low dose group included patients prescribed less than 27 mg a day of methylphenidate (median value), and the high dose group included patients with prescriptions of 27 mg or more.

We conducted several sensitivity analyses in which we examined additional risk periods: a washout period, the day of initiation, and a pre-initiation exposure risk period. The washout period was used to account for the possibility that patients might not take their medicine strictly according to the prescription instructions and might be exposed to drugs for longer than prescribed. Washout periods were defined as 1-3 days, 4-7 days, and 8-14 days after the end date of prescription supply. The day of initiation was also examined separately, and we also included two consecutive 30 day pre-initiation exposure risk periods before the date the drug was first prescribed. This was done to account for the potential that the occurrence of the adverse event could influence exposure—that is, those patients who experience a cardiovascular event might be more or less likely to be started on methylphenidate.

The statistical application program SAS (release 9.4) was used for all statistical analyses. We considered a two tailed value of $P<0.05$ to be significant.

**Results**

We identified and included 1224 eligible participants in the final analysis; 864 had arrhythmias, 396 had disease, 52 had myocardial infarction, 67 had ischaemic stroke, and 44 had heart failure. The mean duration of methylphenidate exposure was 0.5 years for all events except heart failure, which was 0.3 years. The median age at first exposure was 11-13 and median age at first event was 11-13. Boys constituted 75-80% of each of the cohorts. The prevalence of comorbidities varied across adverse events. Depressive episode was the most common comorbidity (n=15, 29% in participants with myocardial infarction), though only nine (13%) participants with ischemic stroke had this condition. In contrast, mental retardation occurred in 12 (18%) participants with an ischemic stroke, while only two (5%) with heart failure had this condition. Antidepressants and antipsychotics were often coprescribed (15-27%). Antiepileptics were used in 17 (25%) with ischemic stroke, though these drugs were used in only 68 (8%) of patients with arrhythmia (table 1).

There was an increased risk of arrhythmia for all risk periods combined (adjusted incidence rate ratio 1.61, 95% confidence interval 1.48 to 1.74), with the highest risk observed on days one to three after the start of treatment (2.01, 1.74 to 2.31). The risk of arrhythmia reduced but remained significantly higher in the subsequent risk periods but was not significantly raised with longer term treatment (>56 days) (table 2). While there was no increased risk of hypertension with methylphenidate exposure overall, when exposure time was divided into several risk periods after initiation, a higher risk was observed on days four to seven after the start of treatment. For myocardial infarction, no increased risk was observed overall, though risk was significantly raised for exposure up to 56 days only. No increased risk was observed with methylphenidate exposure for ischaemic stroke or heart failure. Incidence rates and age adjusted rate ratios are presented in the appendix.

Subgroup analysis for arrhythmia showed that risk was higher in patients with than without congenital heart disease (incidence rate ratio 3.49 (95% confidence interval 2.33 to 5.22) v 1.34 (1.23 to 1.46); table 3).
Table 1 | Characteristics of children and young people with ADHD in study of effect of treatment with methylphenidate

|                        | Arrhythmias | Hypertension | Myocardial infarction | Ischemic stroke | Heart failure |
|------------------------|-------------|--------------|-----------------------|-----------------|--------------|
| No of patients         | 864         | 396          | 52                    | 67              | 44           |
| Median (IQR) duration of exposure (years) | 0.5 (0.1-1.3) | 0.5 (0.1-1.3) | 0.5 (0.1-1.1) | 0.5 (0.2-1.3) | 0.3 (0.1-0.8) |
| Median (IQR) duration of exposure before events (years) | 0.04 (0.0-0.48) | 0.04 (0.0-0.48) | 0.0 (0.0-0.26) | 0.0 (0-0.42) | 0.0 (0-0.13) |
| Median (IQR) duration of exposure after events (years) | 0.20 (0.02-0.75) | 0.17 (0-0.70) | 0.17 (0.04-0.91) | 0.27 (0.04-0.83) | 0.12 (0-0.50) |
| Median (IQR) age at first exposure | 13 (10-15) | 13 (11-15) | 11 (8-16) | 11 (8-14) | 12 (10-16) |
| Median (IQR) age at first outcome | 13 (10-15) | 13 (10-16) | 11 (8-14) | 11 (8-14) | 13 (9-15) |
| No (% of boys)         | 674 (78)    | 317 (80)     | 39 (75)               | 51 (76)         | 33 (75)      |
| No (% of congenital heart disease) | 46 (5.3) | 31 (7.8) | 3 (5.8) | 2 (3.0) | 2 (15.9) |
| No (% with comorbidity* (ICD-10 code): |                          |                          |                          |                          |              |
| Depressive episode (F32-F33, F34.1, F41.2) | 223 (25.8) | 101 (25.5) | 15 (28.9) | 9 (13.4) | 12 (27.3) |
| Tic disorders (F95)    | 76 (8.8)    | 35 (8.8)     | 7 (13.5)              | 6 (9.0)         | 2 (4.6)      |
| Emotional disorders with onset specific to childhood (F93) | 77 (8.9) | 31 (7.8) | 5 (9.6) | 8 (11.9) | 5 (11.4) |
| Conduct disorders (F91) | 82 (9.5) | 34 (8.6) | 1 (1.9) | 3 (4.5) | 3 (6.8) |
| Manic episode (F30)    | 7 (0.8)     | 2 (0.5)      | 0 (0)                 | 0 (0)           | 0 (0)        |
| Bipolar affective disorder (F31) | 44 (5.1) | 15 (3.8) | 2 (3.9) | 3 (4.5) | 1 (2.3) |
| Mental retardation (F70-F79) | 75 (8.7) | 30 (7.7) | 4 (7.7) | 12 (17.9) | 2 (4.6) |
| No (% with comedication† (ATC code): |                          |                          |                          |                          |              |
| Atomoxetine            | 15 (1.7)    | 5 (1.2)      | 2 (2.9)               | 2 (2.9)         | 0 (0.0)      |
| Antipsychotics (ATC: N05A) | 114 (13.2) | 55 (13.9) | 7 (13.5) | 5 (7.5) | 2 (4.6) |
| Antidepressants (ATC: N06A) | 181 (21.0) | 78 (19.7) | 6 (15.4) | 12 (17.9) | 12 (27.3) |
| Antiepileptics (ATC: N03A) | 68 (7.9) | 37 (9.3) | 5 (9.6) | 17 (25.4) | 7 (15.9) |
| Anticholinergic agents (ATC: N04A) | 31 (3.6) | 19 (4.8) | 3 (5.8) | 2 (3.0) | 0 (0) |
| Anxiolytics (ATC: N05B) | 71 (8.2)    | 36 (9.1)     | 0 (0)                 | 4 (6.0)         | 4 (9.1)      |

†Defined as coprescription with methylphenidate during same period.
*Defined as at least one diagnosis or prescription between 1 January 2008 and 31 December 2011.

Table 2 | Risk of cardiovascular adverse events before and after treatment with methylphenidate

| Risk period | Arrhythmias | Hypertension | Myocardial infarction | Ischemic stroke | Heart failure |
|-------------|-------------|--------------|-----------------------|-----------------|--------------|
| No of patients | 630     | 1           | 304                   | 1               | 40           |
| Unexposed RR (95% CI)* | 1.61 (1.48 to 1.76) | 1.07 (0.94 to 1.22) | 1.33 (0.90 to 1.98) | 0.70 (0.49 to 1.01) | 0.54 (0.30 to 0.96) |
| Exposed 234 | 24         | 1           | 10                    | 57              | 40           |
| Time since initiation of methylphenidate (days) |                          |                          |                          |                          |              |
| 1-3  | 24         | 1           | 0                    | 0               | 0            |
| 4-7  | 26         | 1           | 0                    | 0               | 0            |
| 8-14 | 35         | 1           | 0                    | 0               | 0            |
| 15-28| 45         | 1           | 0                    | 0               | 0            |
| 29-56| 51         | 1           | 0                    | 0               | 0            |
| >56  | 53         | 1           | 0                    | 0               | 0            |

*Rate ratio adjusted for age, comorbidity, and co-medication in time varying method.

Table 3 | Subgroup analysis for risk of cardiovascular adverse events before and after methylphenidate according to congenital heart disease and daily dose

| Risk period | Arrhythmias | Hypertension | Myocardial infarction | Ischemic stroke | Heart failure |
|-------------|-------------|--------------|-----------------------|-----------------|--------------|
| Congenital heart disease |                          |                          |                          |                          |              |
| Yes | 46  | 3.49 (2.33 to 5.22) | 31 | 0.95 (0.58 to 1.56) | 3 | NA | 2 | NA | 7 | NA |
| No  | 818 | 1.34 (1.23 to 1.46) | 365 | 0.99 (0.87 to 1.13) | 49 | 1.36 (0.89 to 2.07) | 65 | 0.61 (0.42 to 0.89) | 37 | 0.38 (0.20 to 0.72) |
| Daily dose (mg) |                          |                          |                          |                          |              |
| Low (<27)  | 814 | 1.42 (1.29 to 1.56) | 375 | 1.03 (0.88 to 1.20) | 51 | 1.49 (0.97 to 2.30) | 66 | 0.93 (0.62 to 1.38) | 40 | 0.18 (0.07 to 0.46) |
| High (>27) | 622 | 1.42 (1.29 to 1.55) | 285 | 1.04 (0.91 to 1.19) | 30 | 1.39 (0.88 to 2.20) | 34 | 0.27 (0.15 to 0.49) | 26 | 1.06 (0.56 to 2.02) |

*Rate ratio adjusted for age, comorbidity, and co-medication in time varying method.
†Case series analysis according to daily dose conducted in dataset of patients with prescriptions <27 mg of daily dose, and those with prescriptions ≥27 mg. Patient can be duplicated in both dataset as they might change their dose in study period.
There was no differential effect for hypertensive disease in children with or without congenital heart disease. Numbers of children with congenital heart disease were low for the other outcomes. No differential effects between low and high daily doses of methylphenidate were observed for any outcome.

Sensitivity analysis showed higher risk estimates in the exposed period than obtained in the primary analysis (table 4). The sensitivity analysis for the overall risk of myocardial infarction in exposed time was significant (incidence rate ratio 1.69, 95% confidence interval 1.20 to 2.40), while in the primary analysis the effect was not significant (1.33, 0.90 to 1.98). No increased risk of myocardial infarct was found in the risk period before exposure, while in the washout period the risk was three times higher than in unexposed time.

**Discussion**

**Principal findings**

This study found a significantly increased overall risk of arrhythmia associated with treatment with methylphenidate in children and young people with ADHD. We also identified associations according to the timing of exposure for other cardiovascular adverse events. While the risk of myocardial infarction was not significant overall, we found an increased risk after the first week of treatment, which remained significantly raised for the first two months of continuous treatment. Though there was an increased risk for arrhythmia overall, the risk was substantially higher in patients with existing congenital heart disease. These results are consistent with the biological plausibility that the mechanism of action relates to the effect of methylphenidate on the heart rate. Delayed effects would be expected with myocardial infarction, while more immediate effects would be expected with arrhythmias, as we observed.

**Comparison with other studies**

Previous research has found no evidence of increased risk of stroke or myocardial infarction with methylphenidate, which was consistent with our overall risk. Our time based risk analysis, however, suggests that the risk of myocardial infarction might be higher in the early risk periods (8-14, 15-28, and 29-56 days). This result is of concern and warrants further research. The lack of increased risk for all events with long term treatment might reflect discontinuation of treatment in those who experienced a cardiovascular adverse event or it might be influenced by the limited long term continuous use of methylphenidate in this population.

We found no previous observational studies investigating the risk of heart failure with methylphenidate. There was no increased risk of heart failure in our study, though numbers of events were low. The risk of arrhythmia was increased for all risk periods up to 56 days, and, in particular, risk was highest in patients with congenital heart disease. While the risk of arrhythmia was lower in those without congenital heart disease, a significant risk was still present. The risk of hypertension varied across risk periods and was

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**Table 4. Sensitivity analysis according to definition of risk period before, during, and after treatment with methylphenidate**

| Risk period | Hypertension | Ischemic stroke | Myocardial infarction | Heart failure |
|-------------|--------------|----------------|----------------------|--------------|
| Primary analyses | | | | |
| Unexposed | 6.0 | 3.3 | 1.0 | 1.0 |
| Exposed | 2.3 | 1.3 | 0.4 | 0.4 |
| Secondary analyses | | | | |
| Unexposed | 4.3 | 2.3 | 1.0 | 1.0 |
| Exposed | 2.1 | 1.3 | 0.4 | 0.4 |

**Risk period (days)**

- Pre-risk period (days before treatment with methylphenidate): 0-30
  - Unexposed: 630, RR: 1.61 (1.48 to 1.74)
  - Exposed: 234, RR: 1.07 (0.94 to 1.22)
- Same day as start of methylphenidate treatment: 0-1
  - Unexposed: 252, RR: 1.75 (1.53 to 2.01)
  - Exposed: 92, RR: 1.11 (0.88 to 1.41)
- Time (days) after start of methylphenidate treatment: 1-3
  - Unexposed: 123, RR: 1.75 (1.53 to 2.01)
  - Exposed: 42, RR: 1.11 (0.88 to 1.41)
- Risk period (8-14 days)
  - Unexposed: 123, RR: 1.75 (1.53 to 2.01)
  - Exposed: 42, RR: 1.11 (0.88 to 1.41)
- Washout period (days after treatment): 0-7
  - Unexposed: 123, RR: 1.75 (1.53 to 2.01)
  - Exposed: 42, RR: 1.11 (0.88 to 1.41)

**Notes**

- RR = risk ratio
- 95% CI = 95% confidence interval
- NA = not applicable
- RRR = relative risk ratio
- MRR = multiplicative risk ratio
not consistent across the primary and sensitivity analyses. Some form of detection bias could have influenced the results for hypertension as it is a condition that is non-symptomatic and might have been detected only through routine tests at scheduled follow-up appointments with healthcare providers. We did find a significantly increased risk of hypertension in the four to seven day risk period, which would correspond to the first follow-up visit after the start of treatment. In contrast, all the other cardiovascular events assessed were symptomatic events, the symptoms of which would precipitate attendance independent of a planned visit, hence a more accurate assessment of timing of the event.

We used one of the largest population datasets, and the study design was robust to confounding. A systematic review of previous observational research investigating the risk of cardiovascular adverse events with methylphenidate reported that six out of seven studies in children and adolescents did not show an association. Because of the low number of events, however, many of these previous studies did not have sufficient statistical power, and the study designs used meant that confounding could not be ruled out.8

**Methodological considerations**

To the best of our knowledge, this is the first population based self controlled case series study focusing on the risk of a comprehensive list of cardiovascular adverse events in children exposed to methylphenidate. We also assessed the temporal association between the start of treatment with methylphenidate and cardiovascular adverse events. Previous research has shown that the self controlled case series method produces similar risk estimates to a new users cohort study design and has the advantage of controlling for constant patient specific confounders because of its within person study design.21 Previous studies with cohort designs examined whether children who take methylphenidate have a higher risk of cardiovascular events compared with those who do not. Our study, however, determines whether methylphenidate use "triggers" the occurrence of a cardiovascular adverse event within a patient. Our population included only those children who had the exposure and the event during the study period. As described by Maclure and colleagues, the analysis of a within person study design can be summarized by the expression "why now" rather than "why me" as in a cohort or case-control study.22

We conducted several sensitivity analyses to verify the robustness of our results by applying a washout period, day of initiation, and a pre-initiation exposure risk period. The pre-initiation exposure risk period was used to test the assumption of the self controlled case series design that the occurrence of an event must not alter the probability of subsequent exposures.16,17 We found that the incidence risk ratios were high in the periods before exposure, suggesting that the probability of exposure to methylphenidate is increased after a cardiovascular adverse event for all events except myocardial infarction. By categorizing this period separately from the reference (unexposed) period, we found that the incidence rate ratios in the periods after exposure in the sensitivity analyses were higher than those in our primary analysis (table 4). This is biologically plausible because ADHD itself might be highly correlated with cardiovascular disease. Several observational studies have reported that patients with ADHD have more pre-existing cardiovascular disease than the general population.21,24 Several previous studies applying self controlled case series designs have defined the periods before exposure to control for potential bias.25,27 Our sensitivity analysis suggests that our main analysis might be biased toward the null and might have underestimated the effects of methylphenidate.

Another assumption of the self controlled case series design is that recurrent adverse events must be independent—that is, the occurrence of one event must not alter the probability of a subsequent event occurring.9,17 In our study, once a person has had a cardiovascular adverse event, their short term risk of experiencing another might be increased. Therefore, events might cluster within episodes. We found 10% of children and adolescents had a recurrent diagnosis of cardiovascular adverse events within 30 days. We overcame this limitation, however, by including only the first event.

The final assumption of the self controlled case series design is that the occurrence of the event of interest must not censor or affect the length of the observation period.16,17,21 Our study population covered children and adolescents, and there were only five deaths after the diagnosis of cardiovascular adverse events in 1224 participants. Farrington and colleagues28 and previous studies using a self controlled case series design21,26 have shown that this method might be robust to failure of this assumption. Additionally we found that the main results did not change when we limited analysis to children who survived until the end of the study (data not shown).

**Strengths and limitations**

This study had several strengths. Our study included the entire Korean population and used the claims database for all patients with ADHD collected by the national health insurance program. Until recently, evidence of the cardiovascular effects of drugs for ADHD was limited to a few observational studies. As cardiovascular adverse events are rare in adolescence, these studies had insufficient power to detect rare events. Our results representing a large population in a real world setting suggest that particular cardiovascular adverse events could be associated with exposure, consistent with the mechanism of action of methylphenidate on heart rate. While we did find an increased relative risk, the absolute risk is likely to be low.

Secondly, we performed a self controlled case series study, which controls for patient specific measured and unmeasured confounders that do not vary over time and has previously been shown to produce similar risk estimates to a new user cohort study design.21 While the self controlled case series design adjusts
implicitly for constant patient specific confounders, it is possible that the association observed might be biased because of time varying confounders. Antidepressants, antipsychotics, or antiepileptics are often coprescribed with methylphenidate and could explain some of the association found with cardiovascular adverse events.\(^29\,30\) Though we adjusted all our analyses for time varying comorbidities and comedications, it is possible that other unmeasured time varying confounders could have influenced our results. For example, there might be differences in severity of ADHD among patients, substance use, and precipitating factors that could have influenced both the occurrence of a cardiovascular adverse event and methylphenidate exposure.

The increased risk in the period before treatment suggests that children with cardiovascular events, except myocardial infarction, were started on methylphenidate treatment. This might reflect prescribing in contraindicated patients and is in itself of concern and might indicate that prescribers consider only myocardial infarction to be a contraindication to methylphenidate use. Additionally we found an increased risk of cardiovascular outcomes in the washout period. There are several possible explanations for this. Firstly, the exposure duration might have been underestimated—that is, patients might have still been taking the drug for a longer time than expected because of non-compliance. Secondly, methylphenidate comes in slow release preparations and is metabolized by the liver, either of which could have implications for people who are slow metabolizers, in whom there might be some potential for drug activity beyond the expected supply.

Our findings should be interpreted with caution. Firstly, there is the potential for inaccuracy of coding and incompleteness of records. The outcome measures were limited to patients with diagnoses of cardiovascular adverse events, and we could have missed outcomes not diagnosed. A validation study comparing the diagnosis derived from the health insurance database with the actual diagnosis recorded in patients’ medical records in Korea found that the overall positive predictive value of diagnoses was about 70%.\(^1\) There is no reason to suspect a differential rate of misclassification in exposed and unexposed periods so this is unlikely to introduce bias in our study. Secondly, we defined the study outcome as a primary or secondary diagnosis for cardiovascular adverse events. Though this is likely to have increased the power of our study by capturing all possible cases of cardiovascular events, some minor events might have been captured. Thirdly, given that this is an observational study, non-compliance to the drug prescribed could have influenced our results. If patients did not take their drug after prescription, our exposure measure will be misclassified, which would affect our results. However, there was a median of only one day between repeat prescriptions. This high persistence rate of methylphenidate meant non-compliance was unlikely in our participants. Finally, a safety warning for methylphenidate associated with cardiovascular adverse events was issued in Korea in December 2009. We have assumed that this warning will not have influenced medical practice in our study (2007-11), though it possibly had some influence on the results.

Conclusions
Methylphenidate exposure in children/young people with diagnosis of ADHD is associated with arrhythmia and potentially with myocardial infarction in specific time periods of use. With the increased use of drugs for ADHD globally, the benefits of methylphenidate should be carefully weighed against the potential cardiovascular risks of these drugs in children and adolescents.

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Data sharing: No additional data available.

Transparency: The lead author affirms that the manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

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