Use of interrupted time-series analysis to characterise antibiotic prescription fills across pregnancy: a Norwegian nationwide cohort study

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ABSTRACT

Objectives Antibiotics are the most frequently prescribed medications for pregnant and breastfeeding women. We applied interrupted time-series analysis (ITSA) to describe antibiotic prescription fills patterns in pregnant women and examined recurrent antibiotic fills in subsequent pregnancies.

Designs A population-based drug utilisation study.

Setting Norwegian primary care.

Participants 653,058 pregnancies derived from Medical Birth Registry of Norway linked to the Norwegian Prescription Database (2006–2016).

Main outcome measure Proportion of pregnancies exposed to antibiotics aggregated by week in pregnancy time windows.

Statistical analyses We descriptively analysed antibiotic prescription fills patterns and components in pregnant women. The changes in antibiotic fills in pregnancy time windows were assessed using ITSA. Interruptions points at week 4 to week 7 into pregnancy and delivery were used. Factors associated with antibiotic fills during pregnancy were identified using generalised estimating equations for Poisson regression. Recurrent antibiotic use was estimated using proportion of women who filled antibiotic prescription in a subsequent pregnancy.

Results Antibiotics were filled in 27.6% pregnancies. The ITSA detected an immediate decrease of 0.07 percentage points (95% CI −0.13 to −0.01) in the proportion of exposed pregnancies at 4 weeks after conception, mainly among women taking folate acid before pregnancy. This proportion increased shortly after delivery (immediate change=1.61 percentage points (95% CI 0.31 to 2.91)) then decreased gradually afterwards (change in slope=−0.19 percentage points, 95% CI −0.34 to −0.05)). The strongest factor associated with antibiotic fills during pregnancy was having recurrent urinary tract infections (adjusted OR=2.65, 95% CI 2.59 to 2.72). Women who had filled antibiotics during a pregnancy were up to three times more likely to fill antibiotics in the subsequent pregnancies.

Conclusions ITSA highlighted important impact of pregnancy and delivery on antibiotic fillings. Having antibiotic fills in a pregnancy was associated with recurrent antibiotic fills in subsequent ones.

INTRODUCTION

Antibiotics are the most frequently prescribed medications for pregnant and breastfeeding women ranging from 20% to 49%.1–8 In outpatient setting, the most common indications for prescribing antibiotics during pregnancy are respiratory and urinary tract infections (UTIs). Notably, UTIs are highly prevalent during the second and third trimester (up to 30%).4,5 In the postpartum period, antibiotics are also often prescribed to treat infections related to delivery and breastfeeding (eg, mastitis).5 Several bacterial infections pose a greater risk to mother and child than the antibiotics used to treat the infections. In general, pregnant and breastfeeding women should be treated like other patients; antibiotics should only be prescribed when strictly needed, narrow-spectrum antibiotics should be preferred over broad-spectrum ones, and adherence should be promoted by adequate patient counselling.7–9 Nevertheless, the prescription of antibiotics with uncertain safety profile for these patients can be a tradeoff between treating infections and protecting the mother and child against potential side effects. Safety concerns about these antibiotics are based on either theoretical consideration, preclinical findings or conflicting results from epidemiological studies.10 Examples include fetal tooth discoloration and inhibition of bone growth caused by tetracyclines, ototoxicity by certain
aaminoglycosides, and concern regarding teratogenicity of fluoroquinolones. Such concerns impact clinical recommendations on how individual groups of antibiotics should be used during and after pregnancy. Consequently, antibiotic use during pregnancy and breastfeeding period may differ substantially from the period before pregnancy in term of prescribing rate, type of prescribed antibiotics and indication.

Differences in antibiotic fills throughout pregnancy (eg, higher filling rates in late pregnancy) compared with the periods before and after pregnancy also suggest potential impact of pregnancy-related events such as planning or awareness of pregnancy and delivery on antibiotic fills. Novel techniques of determining and visualising drug utilisation like interrupted time-series analysis (ITSA) using the start of pregnancy or delivery as break point can improve insight on the impact of these events, including both direction and magnitude, on antibiotic prescription fills in pregnant women. Moreover, women with infections and antibiotic use in a given pregnancy may be more prone to recurrent infections and antibiotic use in a subsequent pregnancy. To date, no previous study has follow-up of women from the beginning of their reproductive history to assess antibiotic use in a recurrent pregnancy.

Consistently with other countries, antibiotics are commonly prescribed for pregnant and breastfeeding women in Norway. In addition, clinical guidelines of antibiotic use for pregnant and breastfeeding women in primary care have been developed. Previous study on medication use in Norwegian pregnant women (2005–2015) showed that about 28% of pregnancies in Norway filled antibiotics during their pregnancy in primary care setting. Filling rate during the first trimester (9.5%) was lowest among pregnancy-related periods: prepregnancy (10.2%), second (12.2%) and third (12.9%) trimester, and post partum (16.7%). However, this descriptive study did not explore various components of antibiotic use (eg, broad-spectrum antibiotic use and adherence to guidelines), factors associated with antibiotic fills in pregnant women and potential drivers of the changes in antibiotic fills throughout pregnancy. Having up to date knowledge about patterns of antibiotic utilisation is essential for public health surveillance, for identifying potential areas of improvement in clinical practice and for promoting judicious prescription of antibiotics among pregnant women.

In the present study, we aim at: (1) estimating changes in antibiotic prescription fills in pregnancy time windows using ITSA, (2) identifying components of antibiotic fills and factors associated with antibiotic fills during pregnancy, and (3) describing the rates of recurrent antibiotic use in subsequent pregnancies.

**METHODS**

**Data sources**

We conducted a drug utilisation study based on data from 2006 to 2016 from the Medical Birth Registry of Norway (MBRN) linked to the Norwegian Prescription Database (NorPD) using unique personal identification numbers.

The MBRN is a population-based registry containing information on all births in Norway since 1967. MBRN is based on mandatory notification of all pregnancies lasting more than 12 weeks. In MBRN, the information available for each of pregnancy include maternal identification, demographic information, information on the mother’s health before and during pregnancy, complications during pregnancy and delivery, date of birth and gestational length and other information on the infants.

In brief, the NorPD is a nationwide registry on all prescribed medications irrespective of reimbursement, dispensed at pharmacies to individual patients treated in primary care from 1 January 2004. In NorPD, the information available for each dispensed drug is the trade name, pharmaceutical form, strength, package size, number of packages, reimbursement code and dispensing date. No information on the usage is included. The medications are classified according to the Anatomical Therapeutic Chemical (ATC) classification system.

**Study population**

We included pregnancies having a maternal ID registered in MBRN with valid gestational length (ie, 12 weeks ≤ gestational length ≤ 45 weeks), and child birth year/end of pregnancy between 2006 and 2016 regardless of pregnancy outcomes.

In the analyses of recurrent antibiotic prescription fills in subsequent pregnancies, we restricted to women having their first pregnancy and at least one subsequent pregnancy during the study period.

**Antibiotic prescription fills**

We investigated the prescription fills of antibacterials for systemic use (ATC code starting with J01) registered in NorPD. Prescription fills of topical formulations of antibiotics were not included.

**Pregnancy-related periods**

We defined the following five pregnancy-related periods: one prepregnancy period up to 3 months before pregnancy; three pregnancy periods, including first trimester (days 1–90 after the start of pregnancy), second trimester (days 91–180), and third trimester (day 180 onwards); and one postpartum period up to 3 months after the end of pregnancy. The first day of the last menstrual period (LMP) is estimated by subtracting the gestational age at delivery from the pregnancy end date. Due date, and thus, gestational length and LMP, is estimated by ultrasound, and only if unavailable, by the women’s recall of LMP.

We hypothesised that the recognition of pregnancy preceded changes in antibiotic prescription filling patterns. Consistently with the literature, we assumed early and common pregnancy recognition at fourth and seventh weeks after the start of pregnancy, respectively.
Covariates
We studied maternal sociodemographic factors (including maternal age at delivery, marital status, smoking in pregnancy), factors related to pregnancy or previous pregnancies (including parity, multiple pregnancies, obstetric complications during pregnancy, asthma, recurrent UTI, smoking in pregnancy and previous pregnancy loss), and comedication. Comedication included fertility treatment using clomiphene (ie, ATC code G03GB02) before pregnancy as marker of infertility, use of folic acid before pregnancy as marker of pregnancy planning, medications used for musculoskeletal (ie, ATC M) and nervous system (ie, ATC N)) as these drugs can be used to relieve symptoms associated with infections. Furthermore, drugs from ATC N can be used in patients with psychiatric disorders whose prevalence of infections is higher than the general population. Obstetric complications were measured by obstetric comorbidity index adapted from Bateman et al.19

Measures of antibiotic use
The main outcome measure was the proportion of exposed pregnancies aggregated by week in pregnancy episode (ie, number of exposed pregnancies per 100 pregnancies). Exposed pregnancies were pregnancies with at least one antibiotic prescription filled in a specific pregnancy-related period based on the date of dispensing.

The secondary outcome measures included (1) the proportion of broad-spectrum antibiotic prescriptions and (2) the proportion of first line antibiotic prescriptions among all antibiotic prescriptions in a specific period. Broad-spectrum antibiotics were defined as per European Surveillance of Antimicrobial Consumption Network’s definition.20 First-line antibiotics were defined using clinical guidelines of antibiotic use for pregnant and breastfeeding women in primary care in Norway (online supplemental eTable 1).8

The outcome measure of recurrent use was the proportion of women who filled antibiotic prescription in a subsequent pregnancy.

Data analyses
First, proportion of exposed pregnancies (in overall and by therapeutic classes) and proportion of broad-spectrum antibiotic and first-line antibiotic prescription filled were calculated for different pregnancy-related periods.

Second, we described maternal characteristics and comedications of pregnant women with and without antibiotic prescription fills and explored the extent and patterns of missing data on covariates of interest.

Third, the association between antibiotic prescription filled (dichotomised as yes/no) during pregnancy and the above-mentioned covariates was investigated using generalised estimating equations (GEE) for Poisson regression, limited to pregnancies without missing data on these covariates. This approach made it possible to handle correlated data due to repeated participation of some women in the cohort. Because of explanatory purpose, we included all covariates in the final multivariable model. Adjusted estimates were expressed as ORs with corresponding 95% CIs.

Fourthly, we assessed changes in the proportion of exposed pregnancies aggregated by week using the ITSA with two break points (week 4 and week 7) using data spanning from week 12 before the start of pregnancy to week 42 thereafter. The ITSA was based on the segmented linear regression modelling (detailed modelling strategies in online supplemental appendix).17 21 22 We repeated ITSA with one interruption (end of pregnancy) to assess the changes in proportion of exposed pregnancies aggregated by week following the end of pregnancy (data spanning from week 42 before the end of pregnancy to week 12 afterwards).

Lastly, the proportion of women filling antibiotics, stratified by whether antibiotics were filled in the previous pregnancy, were calculated (results were presented until the fourth pregnancy).

Additional analyses
Sensitivity analysis using normal logistic regression was performed to check the robustness of findings from GEE model.

The use of folic acid before pregnancy is a potential marker of pregnancy planning which may lead to changes in willingness to fill antibiotic prescriptions.23 In Norway, this information is routinely recorded in MBRN. Nearly one-third of general birthing population in Norway report folic acid use prior to pregnancy.24 25 We performed additional ITSA among those with and without acid folic use before pregnancy to assess the potential role of pregnancy planning on antibiotic prescription fills.

We also applied ITSA: (1) for medications used for musculoskeletal and nervous system to contrast the changes detected by ITSA for antibiotics and (2) for main antibiotic therapeutic classes.

Data management and statistical analyses were performed with Stata/MP V.16.0 for Windows.

RESULTS
Antibiotic fills before, during and after pregnancy
The study population consisted of 653 058 pregnancies among 423 604 women having valid maternal ID with gestational length between 12 and 45 weeks ended in 2006–2016 (figure 1).

A total of 180 516 out of 653 058 pregnancies filled at least one antibiotic prescription during pregnancy (27.6%) and the prescription filling rates decreased over the study period (online supplemental eTable 2 and eFigure 1). The proportion of pregnancies exposed to antibiotics in the 3 months before pregnancy, first trimester, second trimester, third trimester and 3 months after pregnancy were 10.9%, 9.9%, 11.3%, 13.2% and 16.1% respectively. Penicillins (J01C) remained the most common antibiotic class filled in any pregnancy periods (>60%) and were increasingly filled during pregnancy and after pregnancy. The top 10 filled antibiotics were...
presented in online supplemental eTable 3. There were more first-line and less broad-spectrum antibiotics filled during pregnancy than before and after pregnancy (online supplemental eTable 4).

Characteristics of antibiotic prescription fills during pregnancy

Table 1 described the characteristics of pregnancies with and without antibiotic prescription fills.

The GEE modelling, performed on 653 046 pregnancies with complete data on all covariates (12 pregnancies excluded because of missing marital status, the only covariate with missing data), revealed increased risk of filling an antibiotic during pregnancy among women with a history of recurrent UTIs (adjusted OR=2.65, 95% CI 2.59 to 2.72), comedication with drugs used for musculoskeletal (adjusted OR=1.55, 95% CI 1.50 to 1.60) and nervous system (adjusted OR=1.83, 95% CI 1.80 to 1.87), low maternal age (adjusted OR=1.39, 95% CI 1.37 to 1.42), and other factors (table 1). Sensitivity analysis using logistic regression yielded consistent findings (results not shown).

Changes in antibiotic filling following the recognition or end of pregnancy

Before the fourth week into pregnancy, the proportion of pregnancies exposed to antibiotic decreased (figure 2A). ITSA showed an immediate change of −0.07 percentage points (95% CI −0.13 to −0.01) in the proportion of exposed pregnancies after this break point while no change in slope was observed. After the seventh week into pregnancy, both level and slope increased but we could not reject the null (online supplemental eTable 5). In the period after the potential recognition of pregnancy (week 4 to week 7 into pregnancy), the proportion of pregnancies exposed to antibiotics gradually increased by 0.01 percentage points/week (95% CI 0.01 to 0.02).

Before the delivery, the proportion of pregnancies exposed to antibiotics remained stable (online supplemental eTable 6 and figure 2B). The proportion increased by 1.61 percentage point (95% CI 0.31 to 2.91) immediately after delivery while the slope decreased remarkably afterwards (change in slope=−0.19 percentage points, 95% CI −0.34 to −0.05) (online supplemental eTable 6).

Additional analyses

Among women using folic acid prior to pregnancy, proportion of pregnancies exposed to antibiotics dropped by 0.11 percentage points (95% CI −0.14 to −0.08) after the 4 weeks into pregnancy breaking point, remained stable between week 4 and week 7 into pregnancy then increased by 0.01 percentage points/week (95% CI 0.01 to 0.02) afterwards (online supplemental eTable 5 and eFigure 2). Among women without prior folic acid use, no major change was detected after the recognition of pregnancy. Yet the proportion increased during the recognition window by 0.02 percentage point/week (95% CI 0.01 to 0.02) (online supplemental eTable 5 and eFigure 2).

The proportion of pregnancies exposed to medications used for musculoskeletal systems dropped by 0.12 percentage points/week during the recognition window (95% CI −0.12 to −0.11). This proportion remained low during the remaining duration of pregnancy (online supplemental eFigure 3). After delivery, the proportion rose steeply (immediate change=0.5 percentage points,
Table 1  Characteristics of pregnancies with and without antibiotic prescription fills and factors associated with antibiotic prescription fills, Norway, 2006–2016, 653 058 pregnancies

| Characteristic                     | Study population | No antibiotic prescription filled | Any antibiotic prescription filled | Generalised estimating equations |
|------------------------------------|------------------|-----------------------------------|-----------------------------------|----------------------------------|
|                                    | N (%)            | N (%)                             | N (%)                             | Univariable OR (95% CI)          | Multivariable Adjusted OR (95% CI) |
| Maternal age                       |                  |                                   |                                   |                                  |                                  |
| ≤24 years                          | 102 308 (15.7)   | 66 916 (14.2)                     | 35 392 (19.6)                     | 1.37 (1.35 to 1.40)              | 1.39 (1.37 to 1.42)              |
| 25–29 years                        | 206 141 (31.6)   | 149 768 (31.7)                    | 56 373 (31.2)                     | 1.00                             | 1.00                             |
| 30–34 years                        | 216 922 (33.2)   | 161 266 (34.1)                    | 55 656 (30.8)                     | 0.93 (0.91 to 0.94)              | 0.89 (0.87 to 0.90)              |
| ≥35 years                          | 127 687 (19.5)   | 94 592 (20.0)                     | 33 095 (18.3)                     | 0.93 (0.92 to 0.95)              | 0.77 (0.76 to 0.79)              |
| Marital status                     |                  |                                   |                                   |                                  |                                  |
| Married/cohabiting                 | 604 916 (92.6)   | 441 287 (93.4)                    | 163 629 (90.6)                    | 1.00                             | 1.00                             |
| Other                              | 48 130 (7.4)     | 31 248 (6.6)                      | 16 882 (9.4)                      | 1.42 (1.39 to 1.48)              | 1.29 (1.26 to 1.31)              |
| Parity                             |                  |                                   |                                   |                                  |                                  |
| 0                                  | 267 432 (42.3)   | 204 627 (43.3)                    | 71 805 (39.8)                     | 1.00                             | 1.00                             |
| 1                                  | 376 626 (57.7)   | 267 915 (56.7)                    | 108 711 (60.2)                    | 1.15 (1.14 to 1.17)              | 1.30 (1.29 to 1.32)              |
| Plurality                          |                  |                                   |                                   |                                  |                                  |
| Singleton                          | 641 972 (98.3)   | 464 500 (98.3)                    | 177 472 (98.3)                    | 1.00                             | 1.00                             |
| Multiple                           | 11 086 (1.7)     | 8 042 (1.7)                       | 3 044 (1.7)                       | 0.99 (0.95 to 1.03)              | 0.93 (0.89 to 0.97)              |
| Obstetric index*                   |                  |                                   |                                   |                                  |                                  |
| 0                                  | 430 615 (65.9)   | 313 376 (66.3)                    | 117 239 (64.9)                    | 1.00                             | 1.00                             |
| 1                                  | 146 744 (22.5)   | 105 435 (22.3)                    | 41 309 (22.9)                     | 1.05 (1.03 to 1.06)              | 1.09 (1.07 to 1.11)              |
| ≥2                                 | 75 699 (11.6)    | 53 731 (11.4)                     | 21 968 (12.2)                     | 1.09 (1.07 to 1.11)              | 1.17 (1.14 to 1.20)              |
| Previous miscarriage or stillbirth |                  |                                   |                                   |                                  |                                  |
| Yes                                | 185 847 (28.5)   | 132 408 (28.0)                    | 53 439 (29.6)                     | 1.07 (1.06 to 1.09)              | 1.05 (1.04 to 1.06)              |
| No                                 | 467 211 (71.5)   | 340 134 (72.0)                    | 127 077 (70.4)                    | 1.00                             | 1.00                             |
| Asthma                             |                  |                                   |                                   |                                  |                                  |
| Yes                                | 30 307 (4.6)     | 19 505 (4.1)                      | 10 802 (6.0)                      | 1.46 (1.42 to 1.49)              | 1.24 (1.21 to 1.28)              |
| No                                 | 622 751 (95.4)   | 453 037 (95.9)                    | 169 714 (94.0)                    | 1.00                             | 1.00                             |
| Recurrent urinary tract infections |                  |                                   |                                   |                                  |                                  |
| Yes                                | 26 448 (4.1)     | 13 212 (2.8)                      | 13 276 (7.4)                      | 2.65 (2.59 to 2.72)              | 2.62 (2.56 to 2.69)              |
| No                                 | 626 570 (95.9)   | 459 330 (97.2)                    | 167 240 (92.6)                    | 1.00                             | 1.00                             |
| Smoking in pregnancy               |                  |                                   |                                   |                                  |                                  |
| Yes                                | 116 510 (17.8)   | 85 406 (18.1)                     | 31 104 (17.2)                     | 0.97 (0.96 to 0.98)              | 0.98 (0.97 to 0.99)              |
| No                                 | 536 548 (82.2)   | 387 136 (81.9)                    | 149 412 (82.8)                    | 1.00                             | 1.00                             |
| Folic acid before pregnancy        |                  |                                   |                                   |                                  |                                  |
| Yes                                | 183 898 (28.2)   | 137 241 (29.0)                    | 46 657 (25.9)                     | 1.00                             | 1.00                             |
| No                                 | 469 160 (71.8)   | 335 301 (71.0)                    | 133 859 (74.1)                    | 1.16 (1.14 to 1.17)              | 1.10 (1.08 to 1.11)              |
| Clomiphene 12 months>pregnancy     |                  |                                   |                                   |                                  |                                  |
| Yes                                | 19 654 (3.0)     | 14 216 (3.0)                      | 5438 (3.0)                        | 0.99 (0.96 to 1.02)              | 1.10 (1.06 to 1.14)              |
| No                                 | 633 404 (97.0)   | 458 326 (96.7)                    | 175 078 (96.7)                    | 1.00                             | 1.00                             |
| Comedication, musculo-skeletal drugs |                |                                   |                                   |                                  |                                  |
| Yes                                | 15 557 (2.4)     | 8888 (1.9)                        | 6669 (3.7)                        | 1.91 (1.85 to 1.97)              | 1.55 (1.50 to 1.60)              |
| No                                 | 637 501 (97.6)   | 463 654 (98.1)                    | 173 847 (96.3)                    | 1.00                             | 1.00                             |
| Comedication, nervous system drugs |                  |                                   |                                   |                                  |                                  |
| Yes                                | 56 414 (8.6)     | 32 719 (6.9)                      | 23 695 (13.1)                     | 1.97 (1.93 to 2.00)              | 1.83 (1.80 to 1.87)              |

Continued
95% CI 0.15 to 0.87) and then dropped to the levels observed before pregnancy. The proportion of pregnancies exposed to medications used for nervous systems followed the similar movement (online supplemental eFigure 3).

Stratified analyses by therapeutics class showed different patterns. Notably, tetracyclines followed the patterns of comedications while the proportion of pregnancies exposed to penicillins increased throughout the pregnancy (online supplemental eFigures 4 and 5).

**Recurrent use of antibiotic in subsequent pregnancies**

The assessment of recurrent antibiotic prescription filling revealed that women who had filled antibiotic prescription in a given pregnancy were 1.7 to 3.1 times more likely to fill an antibiotic prescription again in their subsequent pregnancy (figure 3). Notably, among women who had filled antibiotics in all three first pregnancies, 58.7% filled an antibiotic in their fourth pregnancy compared with 18.9% among women without history of antibiotic prescription fills during pregnancy.

**DISCUSSION**

This study gives an updated nationwide overview of antibiotic prescription fills in pregnant and breastfeeding women in Norway. First, we assessed the antibiotic prescription filling patterns before, during, and after pregnancy and took a step further to determine factors associated with antibiotic prescription fills as well as filling patterns of first-line and broad-spectrum antibiotics. Second, our study confirmed the strong impact of pregnancy, especially pregnancy planning, and delivery.
on proportion of pregnancies exposed to antibiotics and other medications using ITSA—a powerful quasi-experimental method. Third, we brought an attention to a novel aspect of antibiotic utilisation—the recurrent antibiotic fills patterns in pregnant women. These highlighted findings could assist with better dissemination and implementation of future strategies to promote judicious use of antibiotics in pregnancy.

This study also shows the advantages of using modern analytical methods in perinatal drug utilisation studies. ITSA, a modelling strategy to estimate changes following interventions, is widely used to assess impact of policy changes or guidelines on drug utilisation at the national level. The implementation of this method using pregnancy-related events as breakpoints, is particularly useful to visualise and study the effects of pregnancy and delivery on medication filling patterns, yet uptake of this method has been slow in perinatal pharmacoepidemiology. We have been able to identify only one prior study using this method to characterise drug utilisation in pregnancy. This is unfortunate, because of the potential advantages and insights this method might bring to the field.

Over a fourth of pregnancies in Norway filled antibiotic, consistently previous literature. Compared with other nationwide studies estimating the proportion of pregnant women who filled antibiotics among all pregnant women, the filling rate of Norway might be lower than those of Denmark (50.8–37.7%) and UK (34%) but higher than those of Germany (19.7%) and Netherlands (20.8%). The differences in clinical guidelines and prescribing routine between Norway and other countries may explain these variations. Interestingly, the filling rates in pregnant women do not seem to mirror antibiotic consumption in general population with Denmark and Norway sharing the same rates while Netherlands was the lowest consumer.

During pregnancy, the proportion of exposed pregnancies was found to be lowest in the first trimester. This probably reflects the impact of pregnancy recognition because we also detected a sudden drop in the proportion of exposed pregnancies after the period spanning from 4 to 7 weeks into pregnancy using the ITSA. Higher proportion of exposed pregnancies observed later in pregnancy could be explained by screening, detecting and treating asymptomatic bacteriuria and UTIs in pregnancy, as recommended in routine maternity care. This is supported by our finding that having recurrent UTIs was associated with almost a threefold higher likelihood of antibiotic use before pregnancy. The exposure to second-line and third-line antibiotics during pregnancy, which was very limited, could be explained by unrecognised pregnancy or severe infections where the benefits of the antibiotic use outweigh the risks. Importantly, the proportion of pregnancies exposed to tetracyclines, which are contraindicated during the second and third trimester, were less than 0.1%. Besides, the proportion of broad-spectrum antibiotic fills before pregnancy was found to be much lower compared with the rest of Europe and North America. Broad-spectrum antibiotics were even less prescribed during and after pregnancy. This could be explained by a high focus on using narrow-spectrum antibiotics as first-line therapy and low resistance rate to these antibiotics in Norway.

The drop in the proportion of exposed pregnancies detected around the 4–7 weeks into pregnancy (ie, potential pregnancy recognition window) by ITSA could explain the low proportion of exposed pregnancies in the first trimester. Notably, the change was more visible among those who potentially planned their pregnancy (ie, reported folic acid use before pregnancy). However, the impact of pregnancy recognition seems to be modest on antibiotics compared with other medications (including medications used for musculoskeletal and nervous systems, antidepressants and psychostimulants). Notably, after the recognition window, the proportion of pregnancies exposed to antibiotic slowly increased while the proportion of pregnancies exposed to other medications remained low throughout the pregnancy. This gradual increase observed particularly for antibiotics is likely to be driven by the detection of infections during maternity care checkups (first visit recommended between weeks 6 and 10 in Norway) and an increased willingness to prescribe medications after the first trimester (ie, organogenesis). Similarly, the sudden rise detected by ITSA after delivery was the main driver for peak observed in the 3 months after pregnancy. The peak observed shortly after birth could be explained by clinical need to treat or to prevent infections arising from delivery-related wound care and breast feeding. Indeed, the proportion of exposed pregnancies was not high throughout the whole period as seen in the ITSA's visualisation.

Interestingly, we found that women who filled antibiotic in a pregnancy were up to three times more likely to fill antibiotic in a subsequent pregnancy, indicating that antibiotic prescription patterns from one pregnancy is carried over to the next. This could be indicative of a group of women more prone to infections and/or more willing to use antibiotics during pregnancy. Studies have demonstrated that history of previous infections during pregnancy (notably UTIs) highly increased the likelihood of recurrent infections in subsequent
pregnancies. Women successfully treated with antibiotics delivering a healthy child may be more willing to adhere to prescribed antibiotics in a subsequent pregnancy compared with women with no prior experience with antibiotic use in pregnancy.

**Strengths and limitations**

To our knowledge, this is the first study that employs ITSA to examine the effect of pregnancy-related events on antibiotic prescription filling patterns. We believe that this method could be highly beneficial for researchers in perinatal pharmacoepidemiology, and could be applied to all therapeutic areas. Moreover, we are among the first to follow-up women from the beginning of their reproductive history to quantify recurrent antibiotic fills in subsequent pregnancies. This study was conducted on prospectively collected information obtained from linked electronic healthcare registries covering an entire nation. However, this study has some limitations. First, our study did not capture pregnancies lasting less than 12 weeks (spontaneous and induced abortions) as these pregnancies are not recorded in MBRN. The patterns of antibiotic prescription fills of these pregnancies might be different from those of pregnancies included in our study population. Second, our study did not capture antimycotics or other formulations than oral formulations despite that they are often used among women with infections in pregnancy. For example, oral metronidazole which can be prescribed for treating symptomatic bacterial vaginosis was not included as it is not classified under ATC J01. Third, we did not have access to information about indications. Therefore, it is difficult to determine whether a prescription was appropriate or not. Fourth, the filling rate may not truly reflect the prescribing rate. Indeed, in a study based on visits to 458 general practitioners, Fossum et al estimated that 83% of antibiotic prescriptions for pregnant women were filled in the pharmacy. Fifth, we do not know if the filled antibiotic prescriptions were taken by the pregnant women. As a result, the filling rate may overestimate the actual utilisation. Sixth, information regarding breastfeeding is not recorded in MBRN. Breastfeeding mothers may have different antibiotic filling patterns compared with non-breastfeeding ones. Seventh, the ITSA is performed with an assumption that no other event than the interruption could have impact on the outcome measure. For antibiotics, this is far from the reality. Of note, the assumed interruption points at gestational weeks 4 and 7 are serving as estimates rather than cut-points when a pregnancy may be identified. Therefore, the outputs from ITSA must be interpreted with caution. Last but not least, because we included all pregnancies in the ITSA regardless of calendar year of delivery, temporal changes in antibiotic prescription fills during pregnancy were not captured in our study. Overall antibiotic prescription fills (notably macrolides) during pregnancy seems to declining in the recent years, possibly as a result of large-scale effort to improve antibiotic prescribing.

**CONCLUSION**

ITSA approach visualised and highlighted the important impact of pregnancy and delivery on antibiotic prescription filling patterns. This method is a promising analytic tool for perinatal pharmacoepidemiology. Women who filled antibiotic in a pregnancy were more likely to fill antibiotic again in the subsequent pregnancy. This evidence might be helpful for prescribers of antibiotics in pregnancy and healthcare professionals caring for pregnant women in clinical practice.
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