Are women with major depression in pregnancy identifiable in population health data?

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Abstract

Background: Although record linkage of routinely collected health datasets is a valuable research resource, most datasets are established for administrative purposes and not for health outcomes research. In order for meaningful results to be extrapolated to specific populations, the limitations of the data and linkage methodology need to be investigated and clarified. It is the objective of this study to investigate the differences in ascertainment which may arise between a hospital admission dataset and a dispensing claims dataset, using major depression in pregnancy as an example. The safe use of antidepressants in pregnancy is an ongoing issue for clinicians with around 10% of pregnant women suffer from depression. As the birth admission will be the first admission to hospital during their pregnancy for most women, their use of antidepressants, or their depressive condition, may not be revealed to the attending hospital clinicians. This may result in adverse outcomes for the mother and infant.

Methods: Population-based de-identified data were provided from the Western Australian Data Linkage System linking the administrative health records of women with a delivery to related records from the Midwives’ Notification System, the Hospital Morbidity Data System and the national Pharmaceutical Benefits Scheme dataset. The women with depression during their pregnancy were ascertained in two ways: women with dispensing records relating to dispensed antidepressant medicines with an WHO ATC code to the 3rd level, pharmacological subgroup, ‘N06A Antidepressants’; and, women with any hospital admission during pregnancy, including the birth admission, if a comorbidity was recorded relating to depression.

Results: From 2002 to 2005, there were 96698 births in WA. At least one antidepressant was dispensed to 4485 (4.6%) pregnant women. There were 3010 (3.1%) women with a comorbidity related to depression recorded on their delivery admission, or other admission to hospital during pregnancy. There were a total of 7495 pregnancies identified by either set of records. Using data linkage, we determined that these records represented 6596 individual pregnancies. Only 899 pregnancies were found in both groups (13.6% of all cases). 80% of women dispensed an antidepressant did not have depression recorded as a comorbidity on their hospital records. A simple capture-recapture calculation suggests the prevalence of depression in this population of pregnant women to be around 16%.

Conclusion: No single data source is likely to provide a complete health profile for an individual. For women with depression in pregnancy and dispensed antidepressants, the hospital admission data do not adequately capture all cases.

Keywords: Population-based, Data linkage, Pharmacovigilance, Case ascertainment, Depression, Pregnancy, Antidepressant

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Background

Data linkage of administrative data has been a rich resource for Western Australian researchers for a number of years [1-5]. The more recent approval to link national data from the Pharmaceutical Benefits Scheme (PBS) to datasets in the Western Australian Data Linkage System (WADLS) provides new and valuable opportunities to examine birth outcome profiles of prescription medicines dispensed for use during the preconception period and pregnancy. However, most of the datasets were established for administrative purposes and not for health outcomes research. In order for meaningful results to be extrapolated to specific populations, the limitations of the data and linkage methodology need to be investigated and clarified. An important step is to understand the limits of case ascertainment within each dataset.

The Mental Health Services in Australia report uses data from the National Survey of Mental Health and Wellbeing 2007 (N = 8800 Australians aged 16–85 years), and from the National Mental Health Establishments Database of the Australian Institute of Health and Wellbeing [6]. In the foreword to the 2006–2007 report, the director notes, “At this stage of data development, the information we can provide is limited to the number of services, or visits, or prescriptions delivered across Australia. We still have very little information about the number of people involved, or the services used per person. This remains an important data gap that can only be addressed by connecting information for mental health consumers within and across various datasets. This might be achieved by a range of strategies including data linkage as well as the information that might flow from the implementation of e-health” [7]. The report showed the most common type of management reported for mental health-related problems was a medication being prescribed, supplied or recommended by the general practitioner. Antidepressants were the most common medication, followed by anxiolytics, and hypnotics and sedatives.

Depression is a major public health issue in Australia. In 2009–10, more than 10% of all PBS prescription claims were related to mental health conditions [8]. A large Australian study found that around 9% of women experienced depression in the antenatal period and 16% in the postnatal period, [9] so depression, as an identified condition, should be well-represented in the administrative health datasets relating to pregnancy.

It is the objective of this study to investigate the differences in ascertainment between two datasets, using major depression in pregnancy as an example.

Methods

This was a population-based data linkage study investigating pregnancy events in WA from 2002 to 2005. A pregnancy event was defined as a hospital admission record in the Hospital Morbidity Data System (HMDS) with a diagnosis code between O00-O99, based upon the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision, Australian Modification (ICD-10-AM) [10]. De-identified data were provided from the WADLS, linking the records of women with each pregnancy event to any related records in the HMDS, the Midwives’ Notification System (MNS), and the Registry of Births and Deaths. These datasets were linked to each other and to data from the national PBS. The linkages and methodology have been described previously [11,12].

With data linkage, we could overlay the dates of each woman’s pregnancy from the MNS (based upon last menstrual period and delivery date) to the PBS dispenses to each woman within the same time frame, to determine exposures to PBS medicines during pregnancy. Using these dates we also determined the hospital admissions for each woman that occurred during her pregnancy. There were 112 pregnancies without an MNS record and these were excluded in the initial validation of the datasets.

In Australia, community prescriptions (i.e. non-public hospital) are dispensed either as private prescriptions or under one of two subsidisation schemes—the PBS and the Repatriation Pharmaceutical Benefits Scheme. All Australians are eligible to receive subsidised rates for prescribed medicines approved under the PBS, with around 80% of prescriptions dispensed in Australia being subsidised. Patients are grouped into two classes: general and concessional. As the general patient copayment rises, the dispensed prices of many of the cheaper medications fall under this level. In such cases the patient pays the full price and no claim for payment is made under the PBS. For the year ending June 2006, 83.8% of all dispenses and 80.0% of expenditure recorded on the PBS were for concessional patients [13]. New medicines are usually listed on the PBS at the full copayment amount, and hence all dispenses are captured in the data. However, the cost to the patient of older medicines and generic versions tend to fall below the copayment level and so not all medicines have been recorded previously for general patients.

Records from the PBS relating to antidepressant use were ascertained by selecting those dispenses of medicines with a WHO Anatomical Therapeutic Chemical (ATC) code to the 3rd level, pharmacological subgroup, ‘N06A Antidepressants.’ Women with any dispense of a medicine during her pregnancy that included one of these codes were ascertained as a PBS case. The medicines in this group are listed in Table 1. The copayment amount for general patients for fluvoxamine maleate 50 mg tablets and moclobemide 150 mg tablets fell below the subsidy level from April to December 2005 and from
August to December 2005 for fluoxetine hydrochloride 20 mg tablets. Other forms of fluvoxamine maleate and moclobemide were listed at the maximum copayment level during this time. All other antidepressants were listed at the maximum copayment level for the period of the study. The PBS did not collect data on dispenses to patients in public hospitals until December 2004. As depression is a chronic condition, it is not likely that a woman would be dispensed an antidepressant only once and whilst she was in hospital.

The system used in Australia to categorise the risk of drug use in pregnancy is a slightly modified version of the Swedish categorisation (Farmaceutiska Specialiteter i Sverige) and was adopted by the Australian Drug Evaluation Committee (ADEC) in 1989 [14,15]. It includes most of the commonly-used prescription and over-the-counter medicines used in Australia. The categorisations apply only to recommended therapeutic doses in women in the reproductive age group. For pharmaceutical products containing two or more active medicines, the categorisation of the combination is based on the component with the most restrictive categorisation [15]. There are around 950 medicines listed in the ADEC classifications in pregnancy. The antidepressants were classified as B1, B2, B3, C or D for risk for use in pregnancy.

The hospital admission data include a principal diagnosis and up to 20 comorbidities (‘additional diagnoses’) as recorded on the discharge records. The codes are based upon ICD-10-AM [10]. According to the HMDS coding guide that was current during the study, [16] additional diagnoses with the following characteristics need to be coded:

- * require therapeutic treatment;
- * require performance of a diagnostic procedure;
- * increase nursing care and/or monitoring; or,
- * may extend the length of stay in hospital.

‘A condition is not routinely coded just because a patient is on ongoing medication treatment of a condition. However, if the medication is altered or adjusted during the episode of care, the condition should be coded.’

We included any hospital admission during pregnancy as well as the birth admission. The codes we used relating to depression are listed in Table 2. These codes are the same as those suggested in the Private Mental Health Alliance [17] and the National Collaborating Centre for Mental Health [18]. Women with any admission that included one of these codes as a principal diagnosis or comorbidity in the record were ascertained as an HMDS case.

The Australian Bureau of Statistics has released Socio-Economic Indexes for Areas (SEIFA) based on the information collected in the five-yearly Census of Population and Housing. These indexes are widely used measures of

| Table 1 Ascertainment of pregnant women with depression from dispensing data, 2002-2005 |
|-----------------|-----------------|-----------------|-----------------|-----------------|
|                  | Dispenses       | PBS cases       | PBS, no HMDS    | PBS and HMDS    |
| Pregnancy Risk Code | N  | %   | N  | %   | N  | %   | N  | %   |
| Pharmaceutical Benefits Scheme | 20879 | 100.0% | 4485 | 100.0% | 3586 | 80.0% | 899 | 20.0% |
| Sertraline hydrochloride | C | 5536 | 26.5% | 1340 | 29.9% | 1082 | 30.2% | 258 | 28.7% |
| Citalopram hydrobromide | C | 4784 | 22.9% | 1136 | 25.3% | 897 | 25.0% | 239 | 26.6% |
| Paroxetine hydrochloride* | C/D | 3180 | 15.2% | 676 | 15.1% | 565 | 15.8% | 111 | 12.3% |
| Venlafaxine hydrochloride | B2 | 3146 | 15.1% | 581 | 13.0% | 430 | 12.0% | 142 | 15.8% |
| Fluoxetine hydrochloride | C | 1509 | 7.2% | 364 | 8.1% | 280 | 7.8% | 84 | 9.3% |
| Escitalopram oxalate | C | 775 | 3.7% | 241 | 5.4% | 189 | 5.3% | 52 | 5.8% |
| Mirtazapine | B3 | 510 | 2.4% | 140 | 3.1% | 85 | 2.4% | 55 | 6.1% |
| Fluvoxamine maleate | C | 412 | 2.0% | 140 | 3.1% | 115 | 3.2% | 25 | 2.8% |
| Amitriptyline hydrochlorine | C | 273 | 1.3% | 105 | 2.3% | 79 | 2.2% | 26 | 2.9% |
| Dothiepin hydrochloride | C | 376 | 1.8% | 78 | 1.7% | 62 | 1.7% | 16 | 1.8% |
| Moclobemide | B3 | 92 | 0.4% | 40 | 0.9% | 34 | 0.9% | 6 | 0.7% |
| Doxepine hydrochloride | C | 137 | 0.7% | 32 | 0.7% | 20 | 0.6% | 12 | 1.3% |
| Reboxetine mesilate | B1 | 89 | 0.4% | 24 | 0.5% | 16 | 0.4% | 8 | 0.9% |
| Imipramine hydrochloride | C | 31 | 0.1% | 8 | 0.2% | 6 | 0.2% | <5 | 0.2% |
| Mianserin hydrochloride | B2 | 9 | 0.0% | 6 | 0.1% | <5 | 0.1% | <5 | 0.2% |
| Nefazodone hydrochloride | B3 | 14 | 0.1% | 5 | 0.1% | <5 | 0.1% | <5 | 0.1% |
| Nortriptyline hydrochloride | C | 6 | 0.0% | <5 | 0.1% | <5 | 0.1% | 0 | 0.0% |

Pregnancy Risk Code: Australian category of risk for the medicine’s use in pregnancy.
* changed from category C to D, September 2005.
relative socio-economic status at a small geographic area level. The indexes rank and identify areas that are relatively more, or less, disadvantaged. They provide contextual information about the area in which a person lives. The indexes have been obtained by principal components analysis which summarises the information from a variety of social and economic variables, calculating weights that will give the best summary for the underlying variables. The categories of variables include income, education, employment, occupation and housing [19].

Three distinct groups were identified: HMDS cases only, PBS cases only, and cases in both datasets. Within the HMDS cases only, two subsets were found: those women dispensed PBS medicines other than antidepressants, and women not dispensed any PBS medicines. These two subsets of HMDS cases are important and we wanted to distinguish between these HMDS groups. The women who were not dispensed any PBS items may be a healthier group; they may be women who had planned their pregnancy and wished to avoid in utero exposure to prescription medicines; or, they may have only used medicines which were not captured by the PBS collection (for example: antibiotics dispensed to general patients, complementary medicines, over the counter medicines). Comparisons of demographic, pregnancy, labour and delivery characteristics using MNS data were also made between the women identified with depression using the PBS (any PBS cases) and the remaining women identified with depression using the hospital admissions (HMDS only cases). Odds ratios with 95% confidence intervals (OR; 95% CI) were calculated for all comparisons of prevalence. Student’s t-tests were used to compare the means of continuous measures such as maternal age and gestation.

The WADLS uses the Automatch software package [20] with probabilistic matching based upon medical record number, surname, first given name and initial, date of birth, sex and address as the principal matching fields. Missed links have been estimated at 0.11% [21]. The WADLS has been validated previously [21,22] and has been used extensively for health research [23]. All records for this study were also validated internally. For example, sex and dates of birth or death were checked across each source. The researchers received all data in a de-identified form from the WADLS. The datasets were analysed using SAS software, version 9.2 [24]. To fulfil the requirements of ethics committees’ approvals relating to individual privacy, we have not reported cell sizes with less than five study subjects. This project has approval from the Human Research Ethics Committees of The University of Western Australia and the Department of Health WA.

Results

Based upon hospital admission records and midwives’ notifications, there were 96698 birth events in WA from 2002 to 2005.

### Table 2 Ascertainment of pregnant women with depression from admissions data, 2002–2005 by ICD-10 codes - 3010 HMDS cases

| Hospital Morbidity Data System* | HMD cases | HMD cases, no PBS | HMD cases and PBS (AD) | HMD cases and PBS (not AD) |
|--------------------------------|-----------|------------------|------------------------|---------------------------|
| N                              | %         | N                | %                      | N                         | %                      |
| HMDS cases                      | 3010      | 100.0%           | 978                    | 32.5%                     | 899                    | 29.9%                  | 1133                    | 37.6%                    |
| Mental disorders and diseases of the nervous system complicating pregnancy, childbirth and the puerperium | 2484 | 82.5% | 791 | 80.9% | 746 | 83.0% | 947 | 83.6% |
| F32 Depressive episode          | 628       | 20.9%            | 112                    | 11.5%                     | 399                    | 44.4%                  | 117                     | 10.3%                    |
| Personality history of other mental and behavioural disorders | 540 | 17.9% | 190 | 19.4% | 151 | 16.8% | 199 | 17.6% |
| F33 Recurrent depressive disorder | 157       | 5.2%             | 37                     | 3.8%                      | 65                     | 7.2%                   | 55                      | 4.9%                     |
| F41.2 Mixed anxiety and depressive disorder | 97 | 3.2% | 24 | 2.5% | 48 | 5.3% | 25 | 2.2% |
| F31.9 Bipolar affective disorder, unspecified | 48 | 1.6% | 5 | 0.5% | 27 | 3.0% | 16 | 1.4% |
| F31.7 Bipolar affective disorder, currently in remission | 9 | 0.3% | <5 | 0.4% | <5 | 0.1% | <5 | 0.4% |
| F31.3 Bipolar affective disorder, current episode mild or moderate depression | <5 | 0.1% | 0 | 0.0% | <5 | 0.2% | <5 | 0.1% |
| F31.4 Bipolar affective disorder, current episode severe depression without psychotic symptoms | <5 | 0.0% | 0 | 0.0% | <5 | 0.1% | 0 | 0.0% |
| F31.6 Bipolar affective disorder, current episode mixed | <5 | 0.0% | 0 | 0.0% | 0 | 0.0% | <5 | 0.1% |
| F31.8 Other bipolar affective disorders | <5 | 0.1% | 0 | 0.0% | <5 | 0.2% | <5 | 0.2% |
| F31.5 Bipolar affective disorder, current episode severe depression with psychotic symptoms | 0 | 0.0% | 0 | 0.0% | 0 | 0.0% | 0 | 0.0% |

AD: antidepressant.

* totals do not equal the sum of the column as women may be diagnosed with more than one disorder or have more than one admission during their pregnancy.
Ascertainment of cases from dispensing records (PBS cases)
At least one antidepressant was dispensed to 4485 (4.6%) women having a birth event with a total of 20879 dispenses of 17 generic medicines (PBS cases): Table 1. There were 38 different forms of antidepressant dispensed with 24 (63%) dispensed to at least 10 pregnant women. Nearly half of all dispenses of an antidepressant were for sertraline hydrochloride (26.5%) or citalopram hydrobromide (22.9%). During the period of this study, 96.1% of the dispenses of antidepressants under the PBS to the pregnant women were as “Restricted Benefit” use which means the medicine can only be prescribed for specific therapeutic uses. These uses are listed in the PBS as ‘major depressive disorders,’ ‘obsessive compulsive disorder,’ and ‘panic disorder where other treatments have failed or are inappropriate.’ This means the women being dispensed the antidepressants under the PBS were confirmed by their clinician as having the need for the therapeutic use required.

Ascertainment of cases from hospital admission records (HMDS cases)
There were 108088 hospital admissions during pregnancy with 89% of the women having only one admission during their pregnancy and this admission was for the delivery. There were 3010 (3.1%) women with a comorbidity related to depression recorded on their delivery admission, or other admission to hospital during pregnancy (HMDS cases): Table 2. The most common comorbidity codes recorded were ‘O99.3 mental disorders and diseases of the nervous system complicating pregnancy, childbirth and the puerperium’ (82.5%), ‘F32 depressive episode’ (20.9%) and ‘Z86.5 Personal history of other mental and behavioural disorders’ (17.9%).

Comparison of ascertained cases by dataset
There were a total of 7495 pregnancies identified by either set of records. Using data linkage, we could determine that these cases represented 6596 individual pregnancies (6.8% of 96698 pregnancies). There were 3586 pregnancies that were PBS cases but not HMDS cases (54.4% of all cases) and 2111 pregnancies that were HMDS cases and not PBS cases (32.0% of all cases): Figure 1. Only 899 pregnancies were found in both groups (13.6% of all cases). The areas of the circles in the Venn diagram (Figure 1) are proportional to the number of cases ascertained in each dataset. Using a simple capture-recapture analysis, [25] the estimated total number of cases of depression in the 96698 pregnant women would be 15007 (95% CI: 14274–15741), i.e. 15.5%.

Other PBS medicines dispensed
Of the 2111 HMDS cases not dispensed an antidepressant but with depression recorded in their hospital admission records, 978 were not dispensed any medicines under the PBS. The remaining 1133 were dispensed a medicine under the PBS but not an antidepressant. The most frequently dispensed medicines were amoxicillin (N = 290, 25.8%), metoclopramide hydrochloride (N = 200, 17.8%) and cephalaxin (N = 198, 17.6%).

Other morbidities recorded for PBS cases
Of the 4485 cases with an antidepressant dispensed (PBS cases), 899 had a comorbidity of depression recorded, 3481 had other comorbidities recorded that were not related to depression, and 105 did not have any comorbidity recorded on their hospital admissions. The most commonly recorded comorbidities for the 3481 cases are listed in Table 3; with ‘Z72 Problems related to lifestyle’ (N = 727, 20.9%) and ‘O70 Perineal laceration during delivery’ (N = 718, 20.6%) being the most common.

Demographic, pregnancy and delivery characteristics
We investigated the demographic and pregnancy characteristics using data recorded in the MNS: Table 4. We compared the PBS cases (‘any PBS’ cases; N = 4485) with the HMDS cases who were not ascertained in the PBS (‘HMDS only’ cases; N = 2111).

In comparison to the women not dispensed an antidepressant but with a comorbidity record indicating depression, the women dispensed an antidepressant were less likely to have a preterm delivery (0.6; 0.5-0.7), and four times more likely to be Caucasian (4.4; 3.7-5.2). These women were more likely to have had a previous pregnancy (1.3; 1.1-1.4) and a singleton birth (1.6; 1.1-2.2). Their delivery was more likely to be attended by an obstetrician (1.6; 1.4-1.8), more likely to be an elective Caesarean (1.2; 1.1-1.4), and less likely to be an emergency Caesarean (0.7; 0.6-0.8). They were more likely to have a precipitate delivery (1.4; 1.1-1.8) and a local anaesthetic to the perineum (2.0; 1.5-2.6). They were less likely to have their pregnancy complicated by preclampsia (0.6; 0.5-0.8); or to have smoked during their pregnancy (0.8; 0.7-0.9).

Discussion
The objective of this study was to investigate the differences in ascertainment which may arise between a hospital admission dataset and a dispensing claims dataset, using major depression in pregnancy as an example. Using data linkage, we found records for 6596 pregnancies (6.8% all births) that indicated the mother was dispensed an antidepressant during her pregnancy and/or depression was recorded on her hospital admission records. This proportion is similar to a large Australian study around the same period which reported 8.9% women with an antenatal Edinburgh Postnatal Depression Scale >12 and 5.4% >14 [9]. Only 899 pregnancies were found in both groups (13.6% of all cases).
80% of women dispensed an antidepressant did not have depression recorded in their hospital records. 70% of women with depression recorded in their hospital admission record were not dispensed an antidepressant. If the true number of women with antenatal depression is around 15% based upon the capture-recapture algorithm, then there are many pregnant women with undiagnosed or undeclared depression in the administrative health datasets. These results are reflected in one of the statements from the National beyondblue Perinatal Mental Health program: depression and related difficulties affect around 15 per cent of women during pregnancy and early parenthood, and often goes undetected and untreated [9].

Not all women will continue taking antidepressants whilst they are trying to become pregnant or once they discover they are pregnant for fear of fetal harm. These women may be part of the 978 women ascertained in the HMDS without use of any PBS medicines. This group of HMDS cases may also have been dispensed medicines that are not routinely collected in the PBS. The PBS dataset includes only medicines dispensed under subsidy. Some medicines have a wide range of forms of older medicines that are prescribed but no longer fully subsidised so the number of pregnant women identified as treated would be under-estimated.

The women who were dispensed an antidepressant, the PBS cases, were different in many ways from the HMDS only group (Table 4). From the midwives’ data we found these women were more likely to be Caucasian, have already had at least one delivery, and to have a singleton birth. They were more likely to have a higher socio-economic status. Their delivery was less likely to be preterm, to be induced, or to be an emergency Caesarean section. They were more likely to have a precipitate delivery, and a local anaesthetic to perineum. Many of these characteristics suggest a more medically managed pregnancy and possibly better access to medical services.

Table 3 Most frequent comorbidities recorded on hospital admissions, not related to depression, for the 4485 PBS cases

| ICD category | N | % |
|--------------|---|---|
| HMDS comorbidity recorded as depression | 899 | 100.0% |
| no comorbidity recorded | 105 | 100.0% |
| HMDS comorbidity recorded, not related to depression | 3481 | 100.0% |
| Z72 Problems related to lifestyle | 727 | 20.9% |
| O70 Perineal laceration during delivery | 718 | 20.6% |
| O34 Maternal care for known or suspected abnormality of pelvic organs | 608 | 17.5% |
| O99 Other maternal diseases classifiable elsewhere but complicating pregnancy, childbirth and the puerperium | 583 | 16.7% |
| O68 Labour and delivery complicated by fetal distress | 487 | 14.0% |
| O80 Single spontaneous delivery | 400 | 11.5% |
| O72 Postpartum haemorrhage | 360 | 10.3% |
| O62 Abnormalities of forces of labour | 341 | 9.8% |
| O92 Other disorders of breast and lactation associated with childbirth | 339 | 9.7% |
| O36 Maternal care for other known or suspected fetal problems | 288 | 8.3% |
| O32 Maternal care for known or suspected malpresentation of fetus | 276 | 7.9% |
| Z29 Need for other prophylactic measures | 264 | 7.6% |
| O42 Premature rupture of membranes | 262 | 7.5% |
| O69 Labour and delivery complicated by umbilical cord complications | 259 | 7.4% |
| O60 Preterm delivery | 259 | 7.4% |
The very large proportion of women dispensed an antidepressant but without depression recorded on their hospital records (N = 3586/4485; 80%) is of concern. The comorbidities coded for depression need not routinely be coded just because a patient is on ongoing medication. However, if the medication is altered or adjusted during the episode of care, or the patient requires additional monitoring or nursing care, then the condition should be coded [16]. Women with depression that is well-controlled by medication or other therapies may not inform the hospital clinicians of their depression for a range of reasons; [26,27] or, the clinician may judge that their depression did not require adjustment of therapeutic treatment whilst she was admitted. In either situation, the woman may not be recorded in the HMDS as having depression. In primary health care settings, if depression is not routinely asked about, over 50% of cases are missed, highlighting the need for a systematic approach to perinatal psychosocial assessment [9,28] and a similar situation may occur in the hospital setting. Due to the amount of contact that women have with health care providers during pregnancy, this is the best time to start screening, [29] provided that adequately trained staff are available to supply follow-up services to those identified [30].

As more research into the effects of antidepressants on the newborn is published, [31-35] it is hoped that women will advise their hospital care-givers of their use of antidepressants, particularly third trimester use, so that withdrawal symptoms in the newborn may be managed [36]. Since the time period of this study, the National Perinatal Depression Initiative [37] has been...
gathering momentum. The Initiative promotes the provision of routine and universal screening for depression in pregnancy and lactation as clinicians currently use repeatedly to address a variety of hypotheses or public health questions, and the studies are not intrusive. Data linkage provides a very cost-effective approach to postmarketing surveillance and should provide more timely signals of many adverse events in pregnancy than the methods currently in place.

In order for meaningful results to be extrapolated to specific populations, the limitations of the data and linkage methodology need to be investigated and clarified. This study highlights the limits of case ascertainment within each dataset. It also raises concerns around the reporting of depression in hospital records - whether medical staff are aware that pregnant women are taking antidepressants during their pregnancy, and the consequences this may have for the neonate.

Competing interests
The authors declare that they have no competing interests.

Authors' contributions
LC conceived the study, participated in its design and analysis, and drafted the manuscript. LSS, FJS and CB made substantial contributions to acquisition of data, and provided critical review of the manuscript. All authors read and approved the final manuscript.

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