ABSTRACT

Objectives: To describe the effect of in utero exposure to the buprenorphine+naloxone combination product in a rural and remote population.

Setting: A district hospital that services rural and remote, fly-in communities in Northwestern Ontario, Canada.

Participants: A retrospective cohort study was conducted of 855 mother infant dyads between 1 July 2013 and 30 June 2015. Cases included all women who had exposure to buprenorphine+naloxone during pregnancy (n=62). 2 control groups were identified; the first included women with no opioid exposure in pregnancy (n=618) and the second included women with opioid exposure other than buprenorphine+naloxone (n=159). Women were excluded if they had multiple pregnancy or if they were part of a methadone programme (n=16). The majority of women came from Indigenous communities.

Outcomes: The primary outcomes were birth weight, preterm delivery, congenital anomalies and stillbirth. Secondary neonatal outcomes included gestational age at delivery, Apgar scores at 1 and 5 min, NAS Score >7 and treatment for neonatal abstinence syndrome (NAS). Secondary maternal outcomes included the number of caesarean sections, postpartum haemorrhages, out of hospital deliveries and transfer of care to tertiary centres.

Results: No difference was found in the primary outcomes or in the Apgar score and caesarean section rate between in utero buprenorphine+naloxone exposure versus no opioid exposure in pregnancy. Compared to women taking other opioids, women taking buprenorphine+naloxone had higher birthweight babies (p=0.001) and less exposure to marijuana (p<0.001) during pregnancy.

Conclusions: Retrospective data suggest that there likely is no harm from taking buprenorphine+naloxone opioid agonist treatment in pregnancy. Larger, prospective studies are needed to further assess safety.

INTRODUCTION

Opioid dependence in pregnancy is an increasingly common occurrence in rural and remote areas such as Northwestern Ontario, Canada, where up to 28% of pregnancies are exposed to opioid use.1 Our catchment area of 30 000 patients in remote communities who are mostly Indigenous and receive their care to tertiary centres.

bmjopen-2016-011774.

To cite: Jumah NA, Edwards C, Balfour-Boehm J, et al. Observational study of the safety of buprenorphine+naloxone in pregnancy in a rural and remote population. BMJ Open 2016;6:e011774. doi:10.1136/bmjopen-2016-011774

Strengths and limitations of this study

- Opioid misuse is epidemic in rural and remote areas of Northwestern Ontario, Canada, with up to 30% of women exposed during pregnancy. Community-based buprenorphine+naloxone programmes have engaged many rural women in treatment programmes who otherwise would not receive care.
- This is the largest cohort of women exposed to buprenorphine+naloxone in pregnancy and contains detailed information about the daily dose, cumulative dose and exposure time with respect to each trimester of pregnancy.
- While 62 women had exposure to buprenorphine+naloxone in pregnancy, only 3 women had exposure throughout all three trimesters, a further 48 had exposure in the first trimester only and the remainder had variable lengths of exposure.
- Data on illicit substances, smoking and alcohol use during pregnancy were determined by self-report and confirmed with urine drug screens. Data were not collected on other exposures or on the use of other medications such as antidepressants, anxiolytics and folic acid.
- Data were collected retrospectively and were limited to antenatal, maternal and neonatal outcomes. Prospective data and long-term outcomes would provide more robust safety data.
communities in order to provide access to treatment in areas with high rates of opioid dependence and no access to methadone.

The WHO and several national Obstetrics and Gynaecology associations recommend that, when pregnancy is diagnosed, women participating in a buprenorphine+naloxone treatment programme switch to the buprenorphine mono-product because the safety of buprenorphine+naloxone has not been demonstrated in pregnancy. A multicentred randomised controlled trial demonstrated that the buprenorphine mono-product has similar pregnancy outcomes and decreased severity of neonatal abstinence syndrome compared to methadone. Naloxone was added to buprenorphine as a deterrent to illicit use as it precipitates withdrawal from opioids when administered intravenously or intranasally but not via the buccal or sublingual routes. Precipitated withdrawal has been shown to result in adverse pregnancy outcomes.

The caution against using buprenorphine+naloxone in pregnancy is not limited to concerns for withdrawal but also possible teratogenicity. However, to date, there have been no reports of teratogenicity in humans or animals. Congenital anomalies are only one marker of drug safety in pregnancy. The live birth rate, spontaneous abortion rate and stillbirth rates are also markers of safety, and among live births, preterm delivery, low birth weight and functional deficits are factors that may be affected by a medication. Further, the severity of pregnancy outcomes may be modified by the duration and intensity of the exposure to the medication.

In Canada, the buprenorphine mono-product is available only through a special access programme. In our setting, women are counselled to transition from buprenorphine+naloxone to either buprenorphine or long-acting morphine when they present for antenatal care. As a result of delays in obtaining buprenorphine through the special access programme, many women are exposed to buprenorphine+naloxone during early pregnancy and into the second trimester. In addition, for personal reasons, some women opt to remain on buprenorphine+naloxone throughout their pregnancy. This study documents the pregnancy outcomes of a cohort of women from rural and remote communities in Northwestern Ontario who continued to take buprenorphine+naloxone treatment during pregnancy as part of a community-based treatment programme.

METHODS

Participants

Maternal and neonatal data were collected from out-patient antenatal clinic records and inpatient medical records for all pregnancies between 1 July 2010 and 31 July 2015. Cases included all women who had exposure to buprenorphine+naloxone during pregnancy. Two control groups were identified; the first included women with no opioid exposure in pregnancy and the second included women with opioid exposure other than buprenorphine+naloxone. All women who were receiving opioid agonist treatment with buprenorphine+naloxone were advised to switch to the buprenorphine mono-product when available, once pregnancy was diagnosed as per national guidelines. Cases represent those women who elected to stay on buprenorphine+naloxone during their pregnancy. Women were excluded if they had a multiple pregnancy or were taking methadone as part of a treatment programme. All infants room-in with their mother following delivery unless there is a medical or safety reason that precludes rooming-in. The majority of women came from Indigenous communities.

Data collection

A standard case report form was used to collect maternal and neonatal data. The maternal case report form contained information on the health and pregnancy history; smoking, drug and alcohol exposure; and intrapartum data. Smoking was defined by self-reported daily use of cigarettes and was further characterised by the number of cigarettes smoked per day. Alcohol and drug exposure were determined by self-report. Drug exposure was characterised further by urine drug screen results. The neonatal case report form contained information on birth weight, gestational age, Apgars, congenital anomalies and stillbirths.

The primary outcomes for the study was an assessment of the safety of buprenorphine+naloxone, including birth weight, preterm delivery (delivery prior to 37+0 weeks gestational age), congenital anomalies and stillbirth. Secondary neonatal outcomes included gestational age at delivery, Apgar scores at 1 and 5 min, NAS Score >7 and treatment for NAS (two or more NAS Scores that are >7). NAS Scores were calculated using a modified Finnegan Scale at the bedside by nurses who have been trained to use this measure. Secondary maternal outcomes included the number of caesarean sections, postpartum haemorrhages, out of hospital deliveries and transfer of care to tertiary centres.

Statistical analysis

Categorical variables are presented as percentages while continuous variables are presented as a mean with SD. We compared cases to controls using t-test for continuous variables and a Pearson $\chi^2$ test of independence or a Fisher’s Exact test, as appropriate, for categorical data. ORs are presented with the 95% CI. Data analysis was performed with SPSS statistical software V.20 (SPSS, Chicago, Illinois, USA) and Microsoft Excel V.14.1.0 (Microsoft Corp, Redmond, Washington, USA). A p value $<0.05$ indicated statistical significance.

Ethics

Ethics approval was granted by the Sioux Lookout Meno Ya Win Health Centre Research Review and Ethics Committee.
RESULTS

A total of 855 consecutive singleton births were included in the study, of these, 62 had exposure to buprenorphine+naloxone, 618 had no opioid exposure and 159 used illicit opioids during the pregnancy (figure 1). Sixteen women were excluded due to participation in a methadone treatment programme. Twenty-five women were excluded due to a multiple pregnancy. Maternal characteristics are described in table 1. The overall rate of opioid exposure in pregnancy is 27.8%. Data on the racial and ethnic make-up of our study participants as well as data on educational attainment were not collected. Previous studies of this population show that the majority (85%) of women are Indigenous. Educational attainment among Indigenous women living on reserve is low where 57% do not complete high school, 16% receive a high school diploma, 19% participate in post-secondary education and only 4% have a university degree.

A total of 62 women had exposure to buprenorphine+naloxone in pregnancy (see figure 1). Of these women, three women continued prepregnancy buprenorphine+naloxone throughout pregnancy and after delivery. A further 48 women who were taking buprenorphine+naloxone prior to pregnancy switched to buprenorphine alone after the first trimester as per national and international guidelines. Eleven women were induced onto buprenorphine+naloxone during pregnancy, and this occurred in the first trimester for 6 women. The average daily dose of buprenorphine+naloxone was 8.2 ±5.8 mg. Ongoing illicit opioid use was identified in 12 women (19.4%) through routine urine drug screening. Six cases had positive urine drug screens for marijuana. No other illicit substances were identified on urine drug screening. Data were not collected for other psychotropic medications or antidepressant medications.

Cases and controls were of similar age at the time of delivery. However, women who had no opioid exposure in pregnancy had fewer pregnancies and fewer births compared to cases. The OR of Hepatitis C infection for women taking buprenorphine+naloxone compared to no opioid use was OR 15.7, 95% CI 2.6 to 95.6. Women

Figure 1  Patient flow chart.
taking buprenorphine+naloxone were more likely to smoke 1–5 cigarettes per day compared to women in the control groups while women continuing to use illicit opioids were more likely to be heavy smokers (>10 cigarettes per day). There was no difference in alcohol consumption between the three groups.

Prenatal patterns of substance use revealed a high number of women with polysubstance use defined as the use of at least one other illicit substance among women taking buprenorphine+naloxone (12.9%) and those continuing to use illicit opioids (53.5%) (table 2). Treatment with buprenorphine+naloxone decreased the odds of polysubstance use in pregnancy to 0.13 (0.06 to 0.29) compared to women with ongoing illicit opioid use. The most frequent non-opioid drug of abuse was marijuana followed by cocaine (see table 2). Among women continuing to use illicit opioids during pregnancy, the majority (59.7%) used morphine followed by oxycodone derivatives (30.8%). The route of administration for women continuing to use illicit opioids during pregnancy was predominantly intravenous (67.9%) and intranasal (30.8%). Data on the primary opioid of abuse and route of administration were largely missing for women taking buprenorphine+naloxone and as a result a valid comparison cannot be made.

The primary outcome of this study is the safety of in utero exposure to buprenorphine+naloxone (table 3). For the 62 women exposed to buprenorphine+naloxone during pregnancy, the duration of exposure (mean±SD) was 121.4±75.5 days with a daily dose of 8.2±5.8 mg. Among parameters used to define safety of the drug, there was no difference in the birth weight, number of preterm deliveries, number of congenital malformations or number of stillbirths in women taking buprenorphine+naloxone compared to women taking no opioids during pregnancy. Women who continued to use illicit opioids had a statistically significant reduction in birth weight of 262.7 g compared to women treated with buprenorphine+naloxone. There were no stillbirths among the cohort taking buprenorphine+naloxone, five among the women with no opioid exposure and one among the women taking illicit opioids.

There were a total of five congenital malformations in the control group with no opioid exposure and none in the group with illicit opioid exposure. Two infants exposed to buprenorphine+naloxone had congenital malformations: one case of bilateral cleft palate and another case of atrial septal defect. The infant with the bilateral cleft palate had exposure to buprenorphine+naloxone from conception until the third trimester at which point the mother was switched to buprenorphine mono-product. The daily dose ranged from 2 to 4 mg, and the cumulative dose was 525 mg. There was alcohol exposure and smoking (>10 cigarettes per day) during this pregnancy but no other illicit opioids. The infant with the atrial septal defect had exposure to buprenorphine+naloxone from the first trimester until delivery. The daily dose ranged from 2 to 4 mg, and the cumulative dose was 564 mg. There was no other smoking, drug or alcohol exposure during this pregnancy, and there is no family history of congenital heart disease. Data on pre-pregnancy folic acid supplementation and prepregnancy body mass index were not available for either woman. Both women were diagnosed with gestational diabetes.

Secondary neonatal outcomes (table 4) revealed no difference between cases and controls with respect to

### Table 1 Maternal characteristics

| Variable               | A Bup/Nalox (n=62) | B No opioids (n=618) | A–B p Value | C Illicit opioids (n=159) | A–C p Value |
|------------------------|--------------------|----------------------|-------------|--------------------------|-------------|
| Age (years)            | 25.9±4.4           | 25.2±6.3             | 0.21        | 25.5±4.6                 | 0.63        |
| Gravida                | 4.1±2.4            | 3.3±2.3              | 0.009       | 3.9±2.1                  | 0.54        |
| Parity                 | 2.4±1.8            | 1.8±2.0              | 0.03        | 2.2±1.7                  | 0.50        |
| Comorbidities          |                    |                      |             |                          |             |
| Type II diabetes       | 5 (8.1%)           | 26 (4.2%)            | 0.19†       | 3 (1.9%)                 | 0.04†       |
| Gestational diabetes   | 7 (11.3%)          | 63 (10.2%)           | 0.79*       | 14 (8.8%)                | 0.57*       |
| Hypertension           | 6 (9.7%)           | 39 (6.3%)            | 0.29†       | 11 (6.9%)                | 0.58†       |
| HIV                    | 0                  | 0                    |             | 0                        |             |
| Hepatitis B            | 0                  | 0                    |             | 1 (0.6%)                 | >0.99†      |
| Hepatitis C            | 3 (4.8%)           | 2 (0.3%)             | 0.006†      | 12 (7.5%)                | 0.566       |
| Smoking, cig/day       |                    |                      |             |                          |             |
| None                   | 7 (11.3%)          | 327 (52.9%)          | <0.001*     | 20 (12.6%)               | 0.79*       |
| 1–5                    | 33 (53.2%)         | 208 (33.7%)          | 0.002*      | 61 (38.4%)               | 0.045*      |
| 6–10                   | 15 (24.2%)         | 51 (8.3%)            | <0.001*     | 34 (21.4%)               | 0.65*       |
| >10                    | 7 (11.3%)          | 32 (5.2%)            | 0.08†       | 44 (27.7%)               | 0.009†      |
| Alcohol                | 11 (17.7%)         | 134 (21.7%)          | 0.47*       | 41 (25.8%)               | 0.21*       |

Data are n (%) or mean±SD unless otherwise specified.
*χ² test was used.
†When assumptions for the χ² test were not met, the Fisher’s Exact test was used.
gestational age at the time of delivery, Apgar scores, NAS Scores and NAS treatment. Secondary maternal outcomes (Table 4) showed that mothers exposed to buprenor- norphine+naloxone stayed in hospital an extra 1.1 days compared to mothers with no opioid exposure in pregnancy. There was no statistically significant difference in length of stay between cases and illicit opioid using controls. There was no difference in the number of caesar- ean sections, postpartum haemorrhages, out of hospital deliveries or transfers to tertiary care hospitals for cases compared to controls.

DISCUSSION

Almost one-third of the study population were exposed to opioids during pregnancy, but only 5.6% were on opioid agonist treatment prior to pregnancy. There was no evidence of teratogenicity or adverse pregnancy outcomes in a cohort of 62 women exposed to buprenor- norphine+naloxone during pregnancy compared to women who had no opioid exposure during pregnancy. Two cases of congenital malformations were identified in women exposed to buprenorphine+naloxone. One case of bilateral cleft lip and palate included a significant

**Table 2** Prenatal exposure to drugs of abuse

| A Bup/Nalox (n=62) | B No opioids (n=618) | A–B p Value | C Illicit opioids (n=159) | A–C p Value |
|--------------------|---------------------|-------------|-------------------------|-------------|
| **Primary opioid of abuse, n (%)** |
| Morphine 14 (22.6%) | 0 | 95 (59.7%) | <0.001* |
| Oxycodone, Percocet and OxyNeo 5 (8.1%) | 0 | 50 (31.4%) | <0.001† |
| Hydromorphone 4 (6.5%) | 0 | 7 (4.4%) | 0.507† |
| Tylenol+Codeine 0 | 0 | 4 (2.5%) | 0.578† |
| Methadone 0 | 0 | 1 (0.6%) | >0.999† |
| Suboxone 1 (1.6%) | 0 | 9 (5.7%) | 0.289† |
| Unknown 39 (62.9%) | 0 | 7 (4.4%) | <0.001* |
| **Route of opioid administration, n (%)** |
| Intravenous 16 (25.8%) | 0 | 108 (67.9%) | <0.001* |
| Snort/intranasal 5 (8.1%) | 0 | 49 (30.8%) | <0.001† |
| By mouth/per os 0 | 0 | 9 (5.7%) | 0.064† |
| Smoke/inhale 1 (1.6%) | 0 | 7 (4.4%) | 0.447† |
| Not available 42 (67.7%) | 0 | 8 (5.0%) | <0.001* |
| **Frequency of opioid use, n (%)** |
| Occasional (two times per week or less) 1 (1.6%) | 0 | 25 (15.7%) | 0.003† |
| Regular (three times per week or more) 5 (8.1%) | 0 | 26 (16.4%) | 0.111† |
| Daily 11 (17.7%) | 0 | 97 (61.0%) | <0.001* |
| Unknown 45 (72.6%) | 0 | 11 (6.9%) | <0.001* |
| **Other drugs of abuse, n (%)** |
| Polysubstance use 8 (12.9%) | 0 | <0.001† | 85 (53.5%) | <0.001* |
| Marijuana, reported 8 (12.9%) | 60 (9.7%) | 0.424* | 58 (36.5%) | 0.001* |
| Marijuana, positive urine screen 6 (9.7%) | 36 (5.8%) | 0.261† | 44 (27.7) | 0.004† |
| Benzodiazepines 0 | 0 | 3 (1.9%) | 0.561† |
| Cocaine 1 (1.6%) | 0 | 10 (6.3%) | 0.299† |
| Ecstasy 0 | 0 | 1 (0.6%) | >0.999† |

*χ² test was used.  
†When assumptions for the χ² test were not met, the Fisher's Exact test was used.
confiner in the form of in utero alcohol exposure. The second was a case of an atrial septal defect—one of the most common congenital malformations occurring in 1 in 1500 live births. Data regarding other confounders such as folic acid supplementation, exposure to other medications and obesity were not available. In spite of these limitations, the rate of congenital malformations was not significantly different compared to the control group.

Women who continued to use illicit opioids during pregnancy did have babies with a statistically significant lower birth weight compared to those women taking buprenorphine+naloxone. The rate of congenital malformations was not above the rate in the general population although the cohort exposed to buprenorphine+naloxone compared 10 women who took buprenorphine+naloxone in pregnancy to a control group of 31 women who took methadone in pregnancy. The study excluded stillbirth and congenital malformations as well as women who received treatment for NAS during pregnancy rather than the buprenorphine mono-product. In our jurisdiction, the buprenorphine mono-product is available through a special access programme with lengthy delays between applying for an exemption and receipt of the drug, whereas buprenorphine+naloxone is readily available in communities as part of community-based harm reduction programmes. As a result, women are induced onto buprenorphine+naloxone and maintained on this drug until such time that the buprenorphine mono-product becomes available.

Two small retrospective cohort studies are reported in the literature that look at the effect of buprenorphine+naloxone on pregnancy outcomes. The first compared 10 women who took buprenorphine+naloxone in pregnancy to women who took buprenorphine or methadone in six other studies with similar outcome measures. No differences in maternal outcomes or neonatal outcomes were observed with the exception of head circumference which was larger among infants exposed to buprenorphine+naloxone compared to infants who were exposed to methadone withdrawal in utero. The second retrospective cohort study compared 31 women who took buprenorphine+naloxone in pregnancy to a control group of 31 women who took methadone in pregnancy. The study excluded stillbirth and congenital malformations as well as women who received treatment for <30 days prior to delivery. Again, no significant differences in maternal or neonatal outcomes were observed. Infants exposed to methadone were born at an earlier gestational age than buprenorphine+naloxone, but the average gestational age for both was >38 weeks.
Buprenorphine+naloxone is an efficacious treatment for opioid dependence with the advantage of a lower overdose risk and ease of prescribing when compared to methadone.22 Owing to these factors, it is being used widely in opioid treatment programmes in rural and remote settings where methadone treatment is not available.23 Currently women participating in buprenorphine+naloxone treatment programmes who become pregnant are advised to switch to the buprenorphine mono-product out of theoretical concerns of the risk of naloxone to the developing fetus. This study demonstrates no increase rate of congenital malformations, stillbirth or low birth weight for women exposed to buprenorphine+naloxone. Anecdotally women who are stable on buprenorphine+naloxone prior to pregnancy express concern about changing medications and often reluctantly switch to the buprenorphine mono-product on the recommendation of their care provider.

Results from this study support the safety of buprenorphine+naloxone in pregnancy and provide evidence for inclusion of pregnancy as an indication for buprenorphine+naloxone therapy. A policy change of this nature would increase access to care for many women residing in rural and remote settings or where methadone is not an option. Ongoing research and postmarket surveillance would be required to assess rare outcomes. Furthermore, longitudinal studies of the infants exposed to buprenorphine in utero should be performed to assess developmental outcomes. This would add valuable information on the safety of this medication in pregnancy.

Author affiliations
1Northern Ontario School of Medicine, Thunder Bay, Ontario, Canada
2School of Public Health, University of Saskatchewan, Saskatoon, Saskatchewan, Canada
3Anishinaabe Bimaadiziwin Research Program, Sioux Lookout, Ontario, Canada
4Integrated Pregnancy Program, Sioux Lookout Meno Ya Win Health Centre, Sioux Lookout, Canada

Contributors NAJ and CE contributed to data analysis and manuscript preparation. JB-B and KL involved in data collection and provided approval of final draft. JD and LGF contributed to data collection and manuscript preparation. LK involved in data collection, data analysis and manuscript preparation.

Funding This research received no specific grant from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Ethics approval Sioux Lookout Meno Ya Win Health Centre.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement Source data will not be shared publicly as this was not stipulated in the original ethics approval. Data may be obtained by emailing lkelly@mcmaster.ca.

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