Caffeine exposure during pregnancy, small for gestational age birth and neonatal outcome – results from the Norwegian Mother and Child Cohort Study

Dominika Modzelewska 1*, Rino Bellocco 2,3, Anders Elfvin 4, Anne Lise Brantsæter 5, Helle Margrete Meltzer 5, Bo Jacobsson 1,6 and Verena Sengpiel 7

Abstract

Background: Maternal caffeine intake has repeatedly been linked to babies being born small for gestational age (SGA). SGA babies are known to be at increased risk for adverse neonatal outcomes. The aim of this study was to explore the associations between prenatal caffeine exposure and neonatal health.

Methods: The study is based on 67,569 full-term singleton mother-infant pairs from the Norwegian Mother and Child Cohort Study. Caffeine consumption from different sources was self-reported in gestational week 22. Neonatal compound outcomes, namely (1) morbidity/mortality and (2) neonatal intervention, were created based on the Medical Birth Registry of Norway. Adjusted logistic regression was performed.

Results: Caffeine exposure was associated to SGA (OR = 1.16, 95%CI: 1.10; 1.23) and being born SGA was significantly associated with neonatal health (OR = 3.09, 95%CI: 2.54; 3.78 for morbidity/mortality; OR = 3.94, 95%CI: 3.50; 4.45 for intervention). However, prenatal caffeine exposure was neither associated with neonatal morbidity/mortality (OR = 1.01, 95%CI: 0.96; 1.07) nor neonatal intervention (OR = 1.02, 95%CI: 1.00; 1.05 for a 100 mg caffeine intake increase). Results did not change after additional adjustment for SGA status.

Conclusions: Moderate prenatal caffeine exposure (< 200 mg/day) does not seem to impair neonatal health, although prenatal caffeine exposure is associated with the child being born SGA and SGA with neonatal health. We suggest diversity in neonatal outcomes of SGA infants according to the underlying cause of low birth weight.

Keywords: Caffeine, Coffee, Neonatal outcome, Norwegian mother and child cohort study, Pregnancy, Small for gestational age

Background

Caffeine is a plant alkaloid present in different types of beverages and food items such as coffee, tea, soft drinks and chocolate. Excessive caffeine consumption may cause health-related problems [1]. During pregnancy, maternal caffeine clearance is decreased [2], caffeine crosses the placenta easily, and the fetus relies mainly on maternal caffeine clearance. However, there are only few studies on the association of prenatal caffeine exposure and neonatal health. High caffeine intake during pregnancy may result in increased catecholamine levels in the fetus, which may cause placental vasoconstriction [3], and increased fetal heart rate, leading to impaired fetal oxygenation [4]. Some studies reported a higher risk of stillbirth, fetal death [5] or sudden infant death syndrome [6], while others reported no association between caffeine intake and infant death up to 1 year [7].
Our group published a significant association between prenatal caffeine exposure and reduced birth weight as well as increased risk of a baby being small for gestational age (SGA) (odds ratio (OR) = 1.18, 95% confidence interval (CI): 1.10, 1.26) based on 59,123 women from the Norwegian Mother and Child Cohort Study (MoBa) [8]. Consistent findings on the association between caffeine and SGA have been reported in several other studies [9–14].

The term SGA is used as a proxy for intrauterine growth restriction (IUGR). The etiology of IUGR often remains unclear. However, some common fetal, placental and maternal factors associated with IUGR and SGA have been proposed [15, 16]. SGA is strongly associated with neonatal morbidity and mortality [17], and SGA infants are more often admitted to Neonatal Intensive Care Units (NICUs) [18]. In obstetrics, SGA is often used as a surrogate outcome for IUGR and neonatal health as it is widely registered in neonatal records worldwide. However, depending on the underlying cause of reduced birth weight, the risk of neonatal morbidity and mortality might differ. Different underlying SGA causes have been suggested as an explanation for the so-called “birth weight paradox” – neonates born SGA by smokers have a lower infant mortality than neonates born SGA by non-smokers. Hernandez-Diaz proposed that neonates born SGA due to other - more severe SGA causes than smoking such as e.g. congenital malformation have a higher neonatal death risk [19].

Given the known association between caffeine and SGA as well as SGA and neonatal health, we hypothesize that maternal caffeine intake is associated to impaired neonatal health. However, the appropriateness of SGA as a surrogate outcome for IUGR and neonatal health has not yet been established for caffeine exposure studies.

The aim of this study is to assess the association between caffeine exposure from different sources, SGA birth and neonatal health in the Norwegian Mother and Child Cohort Study (MoBa), a large population based study with comprehensive information about lifestyle, maternal health and pregnancy-related conditions.

Methods
Study population and design
This study is based on 67,569 mother-infant pairs from MoBa, an ongoing pregnancy cohort initiated and conducted by the Norwegian Institute of Public Health [20–22]. In short, MoBa is a prospective population-based study. Participants were recruited from all over Norway between 1999 and 2008. Of the invited women, 41% consented to participate. The cohort now includes 114,500 children, 95,200 mothers and 75,200 fathers. Follow-up is conducted by questionnaires at regular intervals and by linkage to national health registries such as the Medical Birth Registry of Norway (MBRN). Follow-up is conducted by questionnaires at regular intervals and by linkage to national health registries. The current study is based on version 8 of the quality-assured data files released for research in 2015 and uses information from the initial questionnaire about general health status and lifestyle filled out around gestational weeks 15 to 17, and the semi-quantitative food frequency questionnaire (FFQ) filled out around gestational week 22. Information from the Medical Birth Registry of Norway (MBRN) is integrated in the MoBa database. MBRN was established in 1967 and contains information about pregnancy, delivery, and health of the mother and the neonate for every live birth, stillbirth or induced abortion after the 12th week of gestation until discharge from the hospital [23]. Hospitals and birth institutions use a standard form to notify MBRN. The MBRN include information from the obstetric record, i.e. data filled in during antenatal visits to a general practitioner, midwife or obstetrician, and information from the medical record for inpatient care, i.e. data recorded from before the time of birth until discharge. The MBRN also contains data from neonatal and paediatric wards on congenital malformations, neonatal diagnoses and procedures performed on infants transferred to those units. The establishment and data collection in MoBa has obtained a license from the Norwegian Data Inspectorate and approval from The Regional Committee for Medical Research Ethics. This cohort is now based on regulations in the Norwegian Health Registry Act. Informed written consent was obtained from each participant. Out of the 114,275 children registered in MoBa, full-term lifeborn singletons without malformations have been selected, for further in- and exclusion criteria see Fig. 1. The current study was approved by the Regional Committee for Medical and Health Research Ethics South East (REK/Sør-Øst 2010/2683).

Caffeine intake
Total daily caffeine intake as well as caffeine intake from various sources (e.g. coffee, tea, soft drinks and chocolate) was estimated based on the self-reported dietary habits in a validated food frequency questionnaire (FFQ) at gestational week 22. Beverage consumption in cups/glasses per day, week or month was reported in specified portion sizes: Coffee (filtered, instant, boiled/pressed, decaffeinated, café latte/cappuccino, espresso or fig/barley) 125 ml/cup, black tea 250 ml/cup, sugar-sweetened or diet cola, soft drinks or chocolate milk 250 ml/glass. Other caffeine sources reported were sandwich spread, desserts, cakes and sweets containing cocoa. Caffeine content was calculated based on FoodCalc [24] and the Norwegian Food Composition Table [25], for detailed information see Sengpiel et al. [8].
Neonatal outcomes
In order to study neonatal outcomes, two composite variables were created: 1) neonatal morbidity or mortality and 2) neonatal intervention. Neonatal morbidity was defined as an Apgar score less than 4 after 5 min or the child being diagnosed with one of the following diagnoses registered in MBRN according to the International Classification of Disease, 10th Edition: birth asphyxia (P21), chronic respiratory disease originating in the perinatal period (P27), intracranial (nontraumatic) hemorrhage of fetus and newborn (P52), meconium ileus/necrotizing enterocolitis (P75, P76, P77, P78.0, P78.1), other disturbances of cerebral status of newborn (P91.0, P91.1, P91.2, P91.6) or retinopathy of prematurity (H35.1). Neonatal mortality was defined as death within 28 days after birth. The selection of the neonatal morbidities was based on the clinical knowledge regarding major conditions occurring among term, preterm and SGA infants. Malformations and genetic syndromes were not included. Neonatal intervention was defined based on data available from the MBRN as a newborn being transferred to the NICU, to receive respiratory or continuous positive airway pressure treatment or treatment with systemic antibiotics (Table 1).

SGA
Information about birth weight and gestational age was obtained from the MBRN. SGA was defined according to Marsal as birthweight below 2 standard deviations (SD) of ultrasound-based growth curves [26]. Since there is no consensus on growth standards, we also ran supplemental analyses defining SGA according to the following approaches:
- Skjaerven, defining SGA as birthweight <10th percentile according to Norwegian population-based growth curves [27],
- Gardosi, defining SGA as birthweight <10th percentile according to customized ultrasound-based growth curves including information on maternal weight and height, parity, and sex of the fetus [28].

Covariates
Information on BMI, maternal smoking status, maternal alcohol consumption and occurrence of nausea, household income, preconceptional folic acid supplementation, and marital status was self-reported in the MoBa

Table 1 The prevalence (number and percentage) of different diagnosis within the neonatal compound outcome variables

| ICD 10 / disorder                                      | N   | %   |
|-------------------------------------------------------|-----|-----|
| Neonatal morbidity / mortality                        | 2124| 100 |
| APGAR score < 4                                       | 119 | 5.6 |
| Birth asphyxia (P21)                                  | 1962| 92.4|
| Other disturbances of cerebral status of newborn (P910–916) | 21 | 1.0 |
| Intracranial (nontraumatic) haemorrhage of fetus and newborn | 96 | 4.5 |
| Retinopathy of the prematurity (H351)                  | 6   | 0.3 |
| Meconium ileus/necrotizing enterocolitis (P75, P76, P77, P78.0, P78.1) | 13 | 0.6 |
| Chronic respiratory disease originating in the perinatal period (P27) | 40 | 1.9 |
| Neonatal death                                        | 57  | 2.7 |
| Neonatal intervention                                 | 9250| 100 |
| Systemic antibiotics                                  | 1447| 15.64|
| Respirator                                            | 133 | 1.44|
| CPAP                                                  | 490 | 5.30|
| NICU admission                                         | 9373| 101.33|

ICD International Classification of Disease, CPAP continuous positive airway pressure, NICU Neonatal intensive care unit. The same child can have different diagnosis or has become subject to several interventions so that the total number is lower than the sum of all diagnosis/interventions.
questionnaires. Information on maternal age at delivery and the baby’s sex was retrieved from the MBRN. The parity variable was based on combined data from MoBa and the MBRN, for detailed information see Sengpiel et al. [3]. Variables were categorized as presented in Table 2.

Statistical analysis
Caffeine intake according to maternal characteristics was analyzed by the Kruskal-Wallis test. The overall association between caffeine intake and neonatal outcomes was analyzed using a logistic regression model, both crude and adjusted for the following categorized variables: maternal pre-pregnancy BMI, household income, maternal education, marital status, parity, maternal age at delivery, smoking status, presence of nausea, folic acid supplementation, planned pregnancy, baby’s sex and total energy intake. Missing data were treated as a category of its own. Additionally, logistic regression analyses were performed including SGA into the model to capture possible indirect effects of prenatal caffeine exposure. All coefficients were reported for a 100 mg change in daily caffeine intake, which equals approximately one cup of coffee. The correlation between covariates was analysed to consider possible collinearity.

Additionally, the assumption of a linear relationship between maternal caffeine intake and log odds of the outcome was assessed by flexible models based on restricted cubic spline regression with five fixed knots.

All statistical analyses were performed using SPSS version 22 and R version 3.3.1.

Results
Caffeine intake
Daily caffeine intake varied between 0 and 1843 mg, with a median value at 58 mg (approximately half a cup of coffee), and 75% of the women consumed less than 123 mg of caffeine per day Fig. 2. Caffeine intake according to maternal characteristics is presented in Table 2. Women who were older, more educated, smoked and had higher incomes consumed more caffeine than those who were younger, non-smoking and had lower incomes.

The main source of caffeine varied across total caffeine intake. Among low caffeine consumers (<100 mg/d), a large portion of caffeine originated from tea, caffeinated soft drinks and chocolate (Fig. 3). With increasing total caffeine intake the fraction of caffeine from coffee increased considerably (Pearson correlation = 0.56, \( P < 0.05 \)).

Caffeine exposure and SGA
The median birth weight in the study population was 3600 g (interquartile range: 3270 g; 3935 g) and 2% of all babies were born SGA. Consistent with our previous study [8], caffeine exposure was associated to SGA (OR = 1.16, 95%CI: 1.10; 1.23). Comparable associations were found if SGA was defined according to Gardosi (14%, OR = 1.1, 95%CI: 1.07; 1.13) or Skjaerven (8%, OR = 1.13, 95%CI: 1.1; 1.17).

SGA and neonatal outcome
The prevalence of the neonatal outcomes is presented in Table 3. As expected, infants born SGA had increased odds for both neonatal morbidity/mortality and neonatal intervention (OR = 3.09, 95%CI: 2.54; 3.78 and OR = 3.94, 95%CI: 3.50; 4.45, respectively). Similar results were found for the other SGA definitions, see Additional file 1).

Prenatal caffeine exposure and neonatal outcomes
Total caffeine intake was not significantly associated with neonatal morbidity/mortality (OR = 1.01, 95%CI: 0.96; 1.07), or neonatal intervention (OR = 1.02, 95% CI: 1.00; 1.05). Additional adjustment for SGA did not affect the estimates (OR = 1.01, 95% CI: 0.95;1.06 for neonatal morbidity/mortality; OR = 1.02, 95% CI: 0.99; 1.05 for neonatal intervention) with similar results for the other SGA definitions, see Additional file 2. Restricting the analysis to never-smokers (n = 61,778) did not change the results (neonatal morbidity/mortality; OR = 1.00, 95% CI: 0.93; 1.06; neonatal intervention OR = 1.02, 95% CI: 0.99; 1.06). Neither did adding interaction terms for maternal BMI, age or the child’s sex change the association (results not shown). No significant association between the different caffeine sources and neonatal outcome variables was found, except for caffeine from chocolate. In the adjusted model, 100 mg increase in chocolate caffeine intake was associated with increased odds for neonatal intervention (OR = 1.59, 95%CI: 1.07; 2.36), see Table 4. There was no evidence of a nonlinear relationship between total caffeine intake (or caffeine intake from different sources) and log odds of neonatal outcomes (\( P > 0.05 \)).

Discussion
The main finding of this study is the lack of any statistically significant association between moderate maternal caffeine intake and neonatal health studied in form of compound neonatal morbidity/mortality and neonatal intervention variables. We previously found a statistically significant association between prenatal caffeine exposure and SGA [8]. Further, we found that SGA was significantly associated with both neonatal outcome variables. While we expected to find a significant association between total caffeine intake and neonatal health, either due to fully (Fig. 4.1) or partly SGA-mediated effects (Fig. 4.2) or caused by residual confounding (Fig. 4.3 or 4.4), maternal caffeine consumption was not associated with neonatal health. Neither did additional adjustment for SGA impact on the association between caffeine exposure and neonatal health. An alternative explanation for our findings would
Table 2 Caffeine Intake According to Maternal Characteristics, n = 67,569, Norwegian Mother and Child Cohort Study, 2002–2009

| Household income, NOK | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|-----------------------|-------|------------------------|-------------------------|----------------------|
| Both partners < 300,000 | 18,703 (28) | 51 (20; 115) | 3 (0; 57) | 6 (0; 29) | < 0.05 |
| One partner ≥ 300,000 | 27,716 (42) | 57 (23; 122) | 6 (0; 66) | 6 (1; 29) | < 0.05 |
| Both partners ≥ 300,000 | 19,263 (29) | 68 (29; 128) | 13 (0; 85) | 6 (1; 40) | < 0.05 |
| Missing | 1887 | 55 (21; 129) | 3 (0; 61) | 6 (0; 29) | < 0.05 |

| Maternal education, years | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|---------------------------|-------|------------------------|-------------------------|----------------------|
| ≤ 12 | 20,840 (32) | 58 (22; 131) | 3 (0; 66) | 3 (0; 17) | < 0.05 |
| 13–16 | 28,136 (43) | 55 (22; 115) | 7 (0; 63) | 6 (0; 29) | < 0.05 |
| 17+ | 17,157 (26) | 68 (28; 124) | 13 (0; 85) | 6 (3; 40) | < 0.05 |
| Missing | 1436 | 60 (22; 127) | 7 (0; 69) | 6 (0; 29) | < 0.05 |

| Marital status | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|----------------|-------|------------------------|-------------------------|----------------------|
| Yes | 64,978 (96) | 58 (23; 122) | 8 (0; 68) | 6 (1; 29) | < 0.05 |
| No | 2591 (4) | 60 (22; 142) | 6 (0; 79) | 3 (0; 17) | < 0.05 |

| Parity | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|--------|-------|------------------------|-------------------------|----------------------|
| 0 | 35,631 (53) | 48 (20; 102) | 6 (0; 48) | 6 (1; 29) | < 0.05 |
| 1 | 20,664 (31) | 68 (28; 137) | 8 (0; 85) | 6 (1; 40) | < 0.05 |
| 2 | 9080 (13) | 85 (33; 175) | 12 (0; 95) | 6 (1; 40) | < 0.05 |
| 3+ | 2150 (3) | 96 (38; 194) | 20 (0; 170) | 6 (0; 40) | < 0.05 |
| Missing | 44 | 50 (19; 130) | 9 (0; 92) | 6 (1; 29) | < 0.05 |

| Maternal age, years | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|---------------------|-------|------------------------|-------------------------|----------------------|
| < 25 | 7712 (11) | 35 (15; 82) | 0 (0; 13) | 3 (0; 17) | < 0.05 |
| 25 to 29 | 22,969 (34) | 48 (20; 102) | 5 (0; 44) | 6 (1; 29) | < 0.05 |
| 30 to 34 | 28,699 (42) | 69 (28; 135) | 12 (0; 85) | 6 (1; 40) | < 0.05 |
| > 34 | 8189 (12) | 92 (40; 180) | 28 (0; 129) | 6 (1; 40) | < 0.05 |

| Alcohol consumption, units/week | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|---------------------------------|-------|------------------------|-------------------------|----------------------|
| No alcohol | 60,155 (89) | 54 (22; 115) | 6 (0; 61) | 6 (1; 29) | < 0.05 |
| < 0.5 | 6214 (9) | 94 (45; 176) | 38 (5; 119) | 17 (3; 40) | < 0.05 |
| > 0.5 | 1199 (2) | 128 (66; 218) | 76 (13; 173) | 17 (3; 40) | < 0.05 |

| Smoking status | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|----------------|-------|------------------------|-------------------------|----------------------|
| Daily | 3596 (5) | 157 (59; 258) | 78 (0; 179) | 3 (0; 17) | < 0.05 |
| Occasionally | 1816 (3) | 110 (45; 201) | 44 (0; 170) | 3 (0; 17) | < 0.05 |
| Never | 61,778 (92) | 55 (22; 113) | 6 (0; 61) | 6 (1; 29) | < 0.05 |
| Missing | 379 | 65 (28; 130) | 12 (0; 75) | 6 (0; 40) | < 0.05 |

| Nausea | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|--------|-------|------------------------|-------------------------|----------------------|
| No | 59,877 (89) | 61 (24; 126) | 8 (0; 79) | 6 (1; 29) | < 0.05 |
| Yes | 7692 (11) | 45 (16; 99) | 0.1 (0; 21) | 6 (0; 29) | < 0.05 |

| Folic acid supplementation | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|---------------------------|-------|------------------------|-------------------------|----------------------|
| No | 37,454 (55) | 62 (24; 131) | 7 (0; 82) | 6 (1; 29) | < 0.05 |
| Yes | 30,115 (45) | 55 (22; 112) | 8 (0; 63) | 6 (1; 29) | < 0.05 |

| Planned pregnancy | N (%) | Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|-------------------|-------|------------------------|-------------------------|----------------------|
| No | 12,994 (19) | 61 (23; 133) | 7 (0; 83) | 6 (0; 29) | < 0.05 |
Table 2  Caffeine Intake According to Maternal Characteristics, n = 67,569, Norwegian Mother and Child Cohort Study, 2002–2009 (Continued)

| Total caffeine, mg/day | Coffee caffeine, mg/day | Tea caffeine, mg/day |
|------------------------|-------------------------|----------------------|
| Yes                    | 54,575 (81)             | 58 (23; 120)         | 8 (0; 67) | 6 (1; 29) |
| Baby’s sex             |                         |                      |          |          |
| Girl                   | 34,340 (51)             | 59 (23; 124)         | 0.13     | 7 (0; 69) | 0.98     | 6 (1; 29) | 0.88 |
| Boy                    | 33,229 (49)             | 58 (23; 122)         | 8 (0; 68) | 6 (1; 29) |
| Quartiles of total energy intake (kcal) | | | | | |
| < 1875                 | 16,882 (25)             | 42 (15; 95)          | < 0.05   | 4 (0; 45) | < 0.05   | 3 (0; 20) | < 0.05 |
| 1875–2222              | 16,883 (25)             | 53 (21; 112)         | 7 (0; 64) | 5 (1; 28) |
| 2223–2651              | 16,927 (25)             | 60 (25; 125)         | 8 (0; 72) | 6 (1; 31) |
| > 2651                 | 16,877 (25)             | 78 (32; 158)         | 10 (0; 89) | 6 (1; 34) |
| BMI                    |                         |                      |          |          |
| < 18.5                 | 2046 (3)                | 56 (22; 114)         | < 0.05   | 7 (0; 64) | < 0.05   | 6 (1; 28) | < 0.05 |
| 18.5 to 24.9           | 43,713 (65)             | 59 (23; 122)         | 8 (0; 74) | 6 (1; 28) |
| 25 to 29.9             | 15,206 (22)             | 59 (23; 126)         | 6 (0; 66) | 6 (1; 28) |
| 30 to 34.9             | 4647 (7)                | 56 (21; 124)         | 3 (0; 48) | 3 (0; 17) |
| 35 to 39.9             | 1358 (2)                | 51 (20; 114)         | 2 (0; 36) | 3 (0; 17) |
| > 40                   | 406 (1)                 | 60 (21; 128)         | 0 (0; 27) | 3 (0; 17) |
| Missing                | 193                     | 70 (25; 137)         | 6 (0; 62) | 6 (0; 40) |

IQR interquartile range, *Kruskal-Wallis test, †Marital Status is defined as either married/cohabiting or not

Fig. 2  Distribution of total daily caffeine intake. Describes the distribution of daily caffeine consumption, median and interquartile range marked in grey, n = 67,569, in the Norwegian Mother and Child Cohort Study 2002 to 2009
be that the effect of SGA on neonatal intervention differs depending on the underlying cause of SGA (Fig. 4.5). Obviously, spurious findings as shown in Fig. 4.3 and 4.4 could be integrated in Fig. 4.5 as well. Assumed there are different types of SGA, one type being statistically – not necessarily causal – linked to maternal caffeine intake and another type due to other – more serious causes for SGA, the neonates being born SGA in association with prenatal caffeine exposure seem to perform better than neonates being born SGA due to other causes. A similar conclusion was presented by Hernandez-Diaz proposing that neonates born SGA due to maternal smoking might have a lower risk of neonatal death than neonates born SGA due to other more severe causes such as congenital malformation [19]. These findings warrant caution when using SGA as a proxy for impaired neonatal health in epidemiologic studies. However, when interpreting the results, it is important to keep in mind that 75% of the study population had caffeine consumption below 123 mg/day, and only 3% of the women had a caffeine intake above 300 mg/day. Therefore, no conclusions can be drawn regarding the effect of high caffeine intake on neonatal health. Aversion to coffee in addition to nausea is a common symptom in early pregnancy resulting in substantially lower caffeine intake in early pregnancy than before pregnancy, and this was also found in our cohort [8].

Results for the different caffeine sources gave the same results except for caffeine from chocolate, which was associated with an increased risk for neonatal intervention in the adjusted analyses. We have no explanation for this and the finding for chocolate is probably due to chance or residual confounding. Habitual intake of coffee and tea is easier to recall than intakes of caffeine-containing soft drinks, chocolate containing foods and plain chocolate as consumption of these items is less regular than coffee and tea. Furthermore, individuals tend to underreport foods

### Table 3: Prevalence of Neonatal Outcomes According to Quartiles of Total Caffeine Intake in the Norwegian Mother and Child Cohort Study, 2002–2009

|                      | Low caffeine intake (<23 mg/day) | Medium caffeine intake (23–58 mg/day) | High caffeine intake (59–123 mg/day) | Very high caffeine intake (>123 mg/day) |
|----------------------|----------------------------------|---------------------------------------|--------------------------------------|----------------------------------------|
| n total              | 16,893                           | 16,892                                | 16,892                               | 16,892                                 |
| Neonatal morbidity/mortality no | 65,445                           | 16,356                                | 16,349                               | 16,362                                 |
| n (%)                | 97                               | 97                                    | 97                                   | 97                                     |
| yes                  | 2124                             | 537                                   | 543                                  | 530                                    |
| n (%)                | 3                                | 3                                     | 3                                    | 3                                      |
| Neonatal intervention no | 58,319                           | 14,559                                | 14,591                               | 14,570                                 |
| n (%)                | 86                               | 86                                    | 86                                   | 86                                     |
| yes                  | 9250                             | 2334                                  | 2301                                 | 2322                                   |
| n (%)                | 14                               | 14                                    | 14                                   | 14                                     |
perceived as unhealthy more than foods perceived as healthy. Estimated intake of caffeine from chocolate was low, 50% of the women had chocolate caffeine intake lower than 4.3 mg/day (interquartile range: 2.2–9.5 mg/day). Thus, the chocolate caffeine variable is less comparable to the other variables and might capture other qualities of a woman’s dietary habits such as having snacks between meals, amount of sweets consumed etc.

To our knowledge, only few studies have evaluated the effect of prenatal caffeine exposure on neonatal health other than birth weight—e.g. neonatal death, sudden infant death and neonatal heart rate. Some studies focused on stillbirth. However, we did not find any studies with comparable outcomes as in this study. A small study from Finland (n = 20) suggests that high caffeine intake during pregnancy (300–600 mg/day, equal to 3–6 cups of coffee/day) may increase catecholamine levels in the fetus and lead to placental vasoconstriction [3]. A study by Resh et al. presented increased contraction rates of surgically isolated fetal hearts when kept in a nutrient solution containing up to 1 mmol/L caffeine. The authors suggested a possible effect of moderate to excessive intake of caffeine during pregnancy on irregular fetal heart rate, leading to impaired fetal oxygenation [4].

A metaanalysis by Chen et al. found an increased risk for pregnancy loss (miscarriage or stillbirth) for maternal caffeine intake above 350 mg/day but not for lower levels [5]. Based on 88,482 women of the Danish National Birth Cohort, Bech et al. published an association between high coffee intake during pregnancy (more than 8 cups/day) and fetal death. However, tea and cola consumption was not associated with fetal death arguing against caffeine being the causal link [29]. In a case-control study from
New Zealand (n = 393 cases and n = 1592 controls), caffeine intake from coffee, tea and cola above 400 mg/day was associated with increased risk for sudden infant death. However, the caffeine intake was assessed retrospectively, opening for recall bias [6]. A prospective Danish cohort study (n = 18,478), found no association between caffeine intake and infant death during the first year of life [7].

Strength and limitations
To our knowledge, with a sample size of 67,569, this study is the largest study performed on the association between caffeine intake and neonatal health. Although the MoBa participation rate is 41%, and the MoBa population differs from the general population of pregnant women by some exposure and outcome characteristics, Nilsen et al. found no differences in eight selected exposure-outcome associations between MoBa participants and the general population registered in the MBRN [22].

The comprehensive MoBa dataset enabled us to control for many important covariates. When studying the potential effects of caffeine exposure, it is crucial to adjust for smoking. Although smoking status was self-reported, the variable has been evaluated against the biological marker cotinine and shown acceptable validity [30]. However, residual confounding cannot be ruled out.

A strength of the MoBa FFQ is that different caffeine sources, portion sizes and preparation methods for coffee could be considered for the estimation of daily caffeine intake. We found similar results for the different caffeine sources studied strengthening the overall finding for prenatal caffeine exposure. Self-reported food consumption might lead to incorrect nutrient intake estimation. However, the MoBa FFQ has been extensively validated based in a sub-population (n = 119), using four-day weighed food diaries and several biomarkers as reference measures. The agreement between the FFQ and the food diaries was particularly high for coffee and tea intake (r = 0.53, r = 0.80 respectively), which are the main sources of caffeine in our study population [31, 32]. Furthermore, the prospective data collection for caffeine intake ensured that a woman’s response was not influenced by her knowledge of the pregnancy outcome.

Neonatal diagnoses were obtained from MBRN, ensuring that diagnoses were registered as given by healthcare professionals. As some of the neonatal conditions might not be diagnosed before the baby leaves the hospital, some of these diagnoses may have been missed. However, the neonatal intervention variable, indicating that the neonate needed medical help in some way, gives complete information for the time until discharge from the hospital.

Supplemental analyses for different common SGA definitions facilitate the interpretation and adaptation of the results in different research settings and healthcare systems. The use of a continuous scale for caffeine intake, as opposed to categorization, ensures that there is no loss of information, power reduction or dilution of the effect of caffeine on neonatal outcome. Categorization risks exposure to misclassification and hampers the comparability among studies. To investigate threshold effects of the exposure, we analysed the possible dose–effect relationship of caffeine intake by restricted cubic spline regression rather than in different categories of caffeine intake.

Conclusion
Moderate maternal caffeine intake is associated with the baby being born SGA, and SGA is strongly associated with impaired neonatal health. However, no significant associations between maternal caffeine intake and neonatal health were found. We suggest that different SGA entities are associated with caffeine intake and neonatal health respectively. Prenatal caffeine exposure seems to be associated with a less severe type of SGA that is not consistently associated to adverse neonatal outcome. These findings warrant caution when using SGA as a proxy for impaired neonatal health in epidemiologic studies. However, when interpreting the results, the relatively low caffeine intake in the cohort, with 75% of the participants consuming less than 123 mg/day, must be kept in mind.

Additional files
Additional file 1: Association Between Small for Gestational Age (SGA) and Neonatal Outcomes, n = 67,569, Norwegian Mother and Child Cohort Study, 2002–2009. aOR – adjusted odds ratios, CI – confidence interval, SGA – small for gestational age. The table shows adjusted odds ratios for the association of small for gestational age (according to different definitions of SGA: Gandosì, Sjøaasen and Marsal) with neonatal outcomes. The Wald 95% confidence intervals are provided for each estimate. ORs are adjusted for: maternal pre-pregnancy body mass index, household income, maternal education, marital status, parity, maternal age at delivery, smoking status, presence of nausea, folic acid supplementation, planned pregnancy and baby’s sex. (DOCX 22 kb)

Additional file 2: Association between caffeine intake and neonatal outcomes, additional adjustment for SGA. CI – confidence interval, OR – odds ratio, SGA – small for gestational age. OR for the outcomes of interest as a function of 100 mg change in total caffeine intake. ORs are adjusted for: maternal pre-pregnancy body mass index, household income, maternal education, marital status, parity, maternal age at delivery, smoking status, presence of nausea, folic acid supplementation, planned pregnancy, baby’s sex, total energy intake, with additional adjustment for small for gestational age (according to a given definition of small for gestational age), n = 67,569, in the Norwegian Mother and Child Cohort Study 2002 to 2009. (DOCX 19 kb)

Abbreviations
BMI: Body mass index; CI: Confidence interval; FFQ: Food frequency questionnaire; IUGR: Intrauterine growth restriction; MBRN: Medical Birth Register of Norway; MoBa: Norwegian Mother and Child Cohort Study; NICU: Neonatal Intensive Care Units; OR: Odds ratio; SGA: Small for gestational age

Acknowledgements
We are grateful to all participating families in Norway who take part in this on-going cohort study.
We want to thank Jonas Bacels1 and Julius Juudak1 for help in the data cleaning process and Margareta Haugen2 for calculation of caffeine intake.

1Department of Obstetrics and Gynecology, Sahlgrenska Academy, Gothenburg University, SE-416 85 Gothenburg, Sweden.
2Norwegian Institute of Public Health, Department of Environmental Exposure and Epidemiology, Domain of Infection Control and Environmental Health, P.O. Box 4404 Nydalen, NO-0403 Oslo, Norway.

Funding
This work was supported by an Erasmus + EU grant received by Dominika Modzelewska.

The Norwegian Mother and Child Cohort Study is supported by the Norwegian Ministry of Health and Care Services and the Ministry of Education and Research, [NIH:NIEHS (contract no N01-ES-75558), NIH/NINDB grant no.1 U01 NS 047537–01 and grant no.2 U01 NS 047537-06A1]. Verena Sengpiel received grants from Goljies minnesfond stiftelsen Sigurd och Elsa [LA2013-0241 “Koffeinintag, födelsevikt och barnutfall!”]. Sthihelsen Fri Mary von Sydows, född Wijk, donationfond (2014 “Koffeinintag, födelsevikt och barnutfall!”) and Wilhelm och Martina Lundgren Vetenskapsfond [1 vetl –119 2014: “Koffeinintag, födelsevikt och barnutfall!”]. Bo Jacobsson received grants by The Swedish Research Council, Stockholm, Sweden (2015–02559), Jane and Dan Olsson Research Foundation, Gothenburg, Sweden as well as funding due to the Agreement concerning research and education of doctors [ALFGBG-426411], Sahlgrenska University Hospital, Sahlgrenska Academy, Gothenburg, Sweden. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Availability of data and materials
The data that support the findings of this study are available from Norwegian Institute of Public Health but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available.

Authors’ contributions
All authors, DM, RB, AE, ALB, HMM, BJ, VS, planned the study, participated in discussing and interpreting results and approved the final manuscript. DM, RB and VS prepared the data, ALB, HMM and VS prepared the exposure variables. DM, AE and VS prepared the outcome variables. DM and VS wrote the first draft.

Ethics approval and consent to participate
The establishment and data collection in MoBa has obtained a license from the Norwegian Data Inspectorate and approval from The Regional Committee for Medical Research Ethics South East (REK/Sør-Øst 2010/2683). Informed written consent was obtained from each participant.

Consent for publication
Not applicable.

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

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1Institute of Clinical Sciences, Department of Obstetrics and Gynecology, University of Gothenburg, Sahlgrenska Academy, SE-416 85 Gothenburg, Sweden.
2Department of Statistics and Quantitative Methods, University of Milano-Bicocca, Milan, Italy.
3Department of Medical Epidemiology and Biostatistics, Karolinska Institutet, Stockholm, Sweden.
4Department of Pediatrics, The Queen Silvia Children’s Hospital, Sahlgrenska University Hospital, Gothenburg, Sweden.
5Department of Environmental Exposure and Epidemiology, Domain of Infection Control and Environmental Health, Norwegian Institute of Public Health, P.O. Box 4404 Nydalen, NO-0403 Oslo, Norway.
6Department of Genetics and Bioinformatics, Domain of Health Data and Digitalisation, Institute of Public Health, Oslo, Norway.
7Department of Obstetrics and Gynecology, Sahlgrenska University Hospital/Ostra, SE-416 85 Gothenburg, Sweden.

Received: 14 September 2018 Accepted: 4 February 2019
Published online: 26 February 2019

References
1. Smith A. Effects of caffeine on human behavior. Food and chemical toxicology: an international journal published for the British Industrial Biological Research Association. 2002;40(9):1243–55.
2. Grosso LM, Bracken MB. Caffeine metabolism, genetics, and perinatal outcomes: a review of exposure assessment considerations during pregnancy. Ann Epidemiol. 2005;15(6):460–6.
3. Kirkinen P, Joupillia P, Koivula A, Vuori J, Puukka M. The effect of caffeine on placental and fetal blood flow in human pregnancy. Am J Obstet Gynecol. 1983;147(8):939–42.
4. Resch BA, Papp JG. Effects of caffeine on the fetal heart. Am J Obstet Gynecol. 1983;146(2):231–2.
5. Chen LW, Wu Y, Neelakantan N, Chong MF, Pan A, van Dam RM. Maternal caffeine intake during pregnancy and risk of pregnancy loss: a categorical and dose-response meta-analysis of prospective studies. Public Health Nutr. 2016;19(7):1233–44.
6. Ford RP, Schluter PJ, Mitchell EA, Taylor BJ, Scagg R, Stewart AW. Heavy caffeine intake in pregnancy and sudden infant death syndrome. New Zealand cot death study group. Arch Dis Child. 1996;78(1):9–13.
7. Widborg K, Kesmodel U, Reich BH, Hedegaard M, Henriksen TB. Maternal consumption of coffee during pregnancy and stillbirth and infant death in first year of life: prospective study. BMJ (Clinical research ed). 2003;326(7386):420.
8. Sengpiel V, Elind E, Bacels J, Nilsson S, Grove J, Myhre R, et al. Maternal caffeine intake during pregnancy is associated with birth weight but not with gestational length: results from a large prospective observational cohort study. BMC Med. 2013;11:42.
9. Bracken MB, Triche EW, Belanger K, Hellenbrand K, Leaderer BP. Association of maternal caffeine consumption with decrements in fetal growth. Am J Epidemiol. 2003;157(5):456–66.
10. Grosso LM, Triche EW, Belanger K, Benowitz NL, Hofvold TR, Bracken MB. Caffeine metabolites in umbilical cord blood, cytchrome P-450 1A2 activity, and intrauterine growth restriction. Am J Epidemiol. 2006;163(11):1035–41.
11. Klebanoff MA, Levine RJ, Clemens JD, Wilkins DG. Maternal serum caffeine metabolites and small-for-gestational age birth. Am J Epidemiol. 2002;155(1):32–7.
12. Orkou J, Henriksen TB, Kesmodel U, Secher NJ. Maternal characteristics and lifestyle factors and the risk of delivering high birth weight infants. Obstet Gynecol. 2003;102(1):115–20.
13. Infante-Rivard C. Caffeine intake and small-for-gestational-age birth: modifying effects of xenobiotic-metabolising genes and smoking. Paediatr Perinat Epidemiol. 2007;21(4):300–9.
14. Vik T, Bakkeiteig LS, Trygg KU, Lund-Larsen K, Jacobsen G. High caffeine consumption in the third trimester of pregnancy: gender-specific effects on fetal growth. Paediatr Perinat Epidemiol. 2003;17(4):324–31.
15. Sharma D, Shastri S, Farahbakhshi N, Sharma P. Intrauterine growth restriction - part 1. The journal of maternal-fetal & neonatal medicine : the official journal of the European Association of Perinatal Medicine, the Federation of Asia and Oceania Perinatal Societies, the International Society of Perinatal Obstet. 2016;1–11.
16. Lee PA, Chernausek SD, Hokken-Koelega AC, Czernichow P. International small for gestational age advisory board consensus development conference statement: management of short children born small for gestational age. April 24-October 1, 2001. Pediatrics. 2003;111(6 Pt 1):1253–1261.
17. Kramer MS. Born too small or too soon. Lancet Glob Health. 2013;1(1):6–7.
18. Hoffmire CA, Chess PR, Ben Saad T, Glantz JC. Elective delivery before 39 weeks: the risk of infant admission to the neonatal intensive care unit. Matern Child Health J. 2012;16(3):1053–62.
19. Hernandez-Diaz S, Schisterman EF, Heman MA. The birth weight "paradox" uncovered? Am J Epidemiol. 2006;164(11):1115–20.
20. Magnus P, Birke C, Vejrup K, Haugan A, Alaker E, Daltveit AK, et al. Cohort profile update: the Norwegian mother and child cohort study (MoBa). Int J Epidemiol. 2016;45(2):382–8.
21. Ronningen KS, Paltiel L, Meltzer HM, Nordhagen R, Lie KK, Hovengen R, et al. The biobank of the Norwegian mother and child cohort study: a resource for the next 100 years. Eur J Epidemiol. 2006;21(8):619–25.
22. Nilsen RM, Vollset SE, Gjessing HK, Skjærven R, Melve KK, Schreuder P, et al. Self-selection and bias in a large prospective pregnancy cohort in Norway. Paediatr Perinat Epidemiol. 2009;23(6):597–608.
23. Irgens LM. The medical birth registry of Norway. Epidemiological research and surveillance throughout 30 years. Acta Obstet Gynecol Scand. 2000;79(6):435–9.
24. Lauritzen J. FoodCalc. [http://www.ibt.ku.dk/jesper/FoodCalc/Default.html].
25. Oslo: NFSANoHDoN-Uo. Matvaretabellen (The Norwegian Food Table, in Norwegian).
26. Marsal K, Persson PH, Larsen T, Lilja H, Selbing A, Sultan B. Intrauterine growth curves based on ultrasonically estimated foetal weights. Acta paediatrica (Oslo, Norway : 1992). 1996;85(7):843–8.
27. Skjærven R, Gjessing HK, Bakketeig LS. Birthweight by gestational age in Norway. Acta Obstet Gynecol Scand. 2000;79(6):440–9.
28. Gardosi J. New definition of small for gestational age based on fetal growth potential. Horm Res. 2006;65(Suppl 3):15–8.
29. Bech HJ, Nohr EA, Vaeth M, Henshaw TB, Olsen J. Coffee and fetal death: a cohort study with prospective data. Am J Epidemiol. 2005;162(10):983–90.
30. Kvalvik LG, Nilsen RM, Skjærven R, Vollset SE, Midttun O, Ueland PM, et al. Self-reported smoking status and plasma cotinine concentrations among pregnant women in the Norwegian mother and child cohort study. Pediatr Res. 2012;72(1):101–7.
31. Brantsæther AL, Haugen M, Alexander J, Meltzer HM. Validity of a new food frequency questionnaire for pregnant women in the Norwegian mother and child cohort study (MoBa). Maternal & child nutrition. 2008;4(1):28–43.
32. Brantsæther AL, Haugen M, Rasmussen SE, Alexander J, Samuelsen SO, Meltzer HM. Urine flavonoids and plasma carotenoids in the validation of fruit, vegetable and tea intake during pregnancy in the Norwegian mother and child cohort study (MoBa). Public Health Nutr. 2007;10(8):838–47.