Prophylactic antibiotics for manual removal of retained placenta during vaginal birth: a systematic review of observational studies and meta-analysis

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Abstract

Background: Manual removal of the placenta is an invasive obstetric procedure commonly used for the management of retained placenta. However, it is unclear whether antibiotic prophylaxis is beneficial in preventing infectious morbidity. We conducted a systematic review to determine the efficacy and safety of routine use of antibiotics for preventing adverse maternal outcomes related to manual placenta removal following vaginal birth.

Methods: A detailed search of MEDLINE, EMBASE, Cochrane library and the CINAHL databases was conducted for non-randomized studies involving women undergoing manual placenta delivery after vaginal births and where antibiotic prophylaxis use was compared with no treatment or placebo to prevent maternal infection. Search terms including ‘delivery, obstetric’, ‘placenta, retained’, ‘anti-infective agents’, and ‘chemoprevention’ were used.

Results: Of the 407 citations that resulted after elimination of duplicates, 81 full texts were potentially eligible after independent assessment of the title and abstracts. Independent review of the full texts identified three eligible cohort studies which were retrospective in design. These studies contained data on two of the pre-specified outcomes, endometritis and puerperal fever. Other secondary outcomes such as perineal infection and/or any infection, hospital stay duration, sepsis, hemorrhage >1000 ml or hospital readmissions were not reported on excluding puerperal fever.

A meta-analysis showed no significant reduction in the incidence of endometritis (odds ratio [OR] 0.84, 95% confidence interval [CI] 0.38 to 1.85, three studies, 567 women) and puerperal fever (OR 0.99, 95% CI 0.38 to 2.27, one study, 302 women).

Conclusions: There is currently no evidence to suggest beneficial effects for routine antibiotic use in women undergoing manual placental removal following vaginal birth. In appropriate settings, further research is required to determine whether a policy of routine antibiotic prophylaxis for the procedure should be maintained or discouraged.

Keywords: Antibiotic prophylaxis, Retained placenta, Vaginal birth, Systematic review, Manual removal, Meta-analysis
Background
In the third stage of labor, after delivery of the infant, the placenta spontaneously detaches from the myometrium [1, 2]. When this does not occur, the placenta is said to be ‘retained’. Studies have shown underlying placental and/or uterine abnormalities to be risk factors for a retained placenta [3]. The time frame for diagnosis post-delivery is still ambiguous, however, a 30–60 min time lapse is widely accepted [4, 5]. Existing reviews reported a varying incidence of 1.5–2.7 % in low-resource and/or high-resource settings respectively, using a 30-min mark-off point [5, 6].

Manual removal of the placenta is indicated if controlled cord traction and the use of uterotonics fails [6–8]. This procedure involves insertion of the hand into the uterus with the aim of separating the placenta from the implantation site, and therefore carries a possible risk of contamination in the uterine cavity. Antibiotic prophylaxis, usually broad spectrum, is routinely administered to reduce infectious morbidities and/or mortalities [9–12]. No evidence exists from randomized control studies or systematic reviews supporting or refuting the practice.

A Cochrane review on the subject initially conducted in 2006 and updated in 2014 did not identify any eligible randomized controlled study [13]. Synthesizing evidence from non-randomized studies is justified in the absence of randomized studies and has been shown to corroborate results from randomized studies regardless of the subject [14]. Hence, we conducted a systematic review on available data from relevant non-randomized studies to determine the efficacy of routine prophylactic antibiotics and if efficacious, the optimal antibiotic regimen for the procedure. This study was conceived as part of the preparation of the evidence base for the World Health Organization (WHO) recommendations for prevention and treatment of maternal peripartum infections.

Methods
Search strategy
Based on a pre-specified protocol prepared in line with guidelines in the Cochrane Handbook for Systematic Reviews [14], we conducted a detailed search on January 28, 2015 for eligible studies on MEDLINE, EMBASE, the Cochrane Library and CINAHL databases using specific search terms that included ‘delivery’, ‘obstetric’, ‘placenta’, ‘retained’, ‘anti-infective agents’ and ‘chemoprevention’ (see Appendix S1).

Initially, studies were selected if they were conducted to answer either of these two questions: (1) What are the effects of routine antibiotic prophylaxis on maternal infectious morbidities and mortality, when used for manual removal of the placenta in vaginal deliveries?; and, (2) What is the comparative effectiveness and safety of different antibiotic regimens used for preventing infectious maternal morbidities during manual removal of the placenta? This systematic review was conducted in accordance with the principles of Declaration of Helsinki. Due to the study design, there was no need for ethics approval as the studies were freely available in the public domain.

Eligibility criteria
All non-randomized studies involving women undergoing manual placental removal after vaginal birth, where the use of antibiotics was compared with no treatment or placebo for prophylaxis against maternal infection. Cluster, quasi-randomized control studies, controlled-before-after studies, cohort and case-control studies were all eligible for inclusion. Comparative studies that reported on comparative use of antibiotic prophylaxis for vaginal births were included while similar studies addressing same comparison but in operative deliveries or a mixed population of operative and vaginal deliveries were excluded, as were studies that contained no data on individual delivery methods.

Data collection and assessment
For this systematic review, the Preferred Reporting Items for Systematic Reviews and Meta-Analyses [15] (PRISMA) method of reporting was used.

The titles and abstracts of all resulting citations from search strategy were examined independently for eligibility by two review authors (CEC and AJQP) irrespective of the publication status (published and unpublished) or language of publication.

Table 1 Characteristics of included studies

| Author, year         | Study design            | Total study population | N (treatment, control) | Enrollment period | Location | Type of obstetric procedure                                      | Available outcomes                  |
|----------------------|-------------------------|------------------------|------------------------|-------------------|----------|------------------------------------------------------------------|-------------------------------------|
| Katsulov 1983 [21]   | Retrospective cohort study | 100                    | 30/70                  | 1976–1977         | Bulgaria | Women who underwent manual placenta removal                     | Endometritis                        |
| Von Rechlin et al. 1988 [22] | Retrospective cohort study | 302                    | 65/237                 | An undefined period of 6 years | Germany | Women undergoing manual placental removal                        | Endometritis, fever greater than 37.5 °C |
| Tandberg et al. 1999 [23] | Retrospective cohort study | 24,750 births          | 61/104                 | 1990–1994         | Norway   | Women who underwent a manual placenta removal                    | Endometritis                        |
After the removal of duplicates, selected studies were then retrieved and further evaluated for inclusion on the basis of full-text appraisal. Disagreements were resolved by discussions with the third review author (EO). Data extraction using a standardized form was performed independently by two review authors (CEC and AJQP), which included information on the study design, facilities, participants, settings and outcomes.

Pre-specified outcomes included infectious morbidities common to obstetric procedures. [5, 16]. Primary outcome was postpartum endometritis as defined by the authors and secondary outcomes included puerperal fever defined as temperature of 38.0 °C or higher, perineal infections, hospital stay duration, sepsis, postpartum hemorrhage, hospital readmission, drug side effects, and neonatal-related outcomes (jaundice, sepsis, intensive care unit admission).

Rating the methodological quality of eligible studies was performed independently by two review authors (CEC and AJQP) using the Newcastle Ottawa scale [17], which involved assessing the methods of participant selection, comparability and outcomes among eligible studies. The quality of the evidence obtained from included studies were further assessed using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach [18–20].

Data analysis/synthesis
Data from included studies (Table 1) was pooled and meta-analysis conducted using a random effects model. Meta-analysis was performed using Revman 5.3 according to the meta-analysis of observational studies in epidemiology (MOOSE) guidelines.

A Mantel Haenszel model for sparse dichotomous data was applied, and adjusted using the random-effects model and presented as odds ratios with a 95 % confidence interval (CI), a p-value of 0.05 and heterogeneity estimates of Tau² and I².

Results
The titles and abstracts of 407 studies that were identified from the keyword search of relevant databases after
elimination of duplicates were independently assessed by two review authors (CEC and AJQP). After initial screening, the full texts of 81 potentially eligible studies were retrieved for further appraisal (Fig. 1). A sizeable number of studies on the subject were published in English and other foreign languages, however, these studies merged data for both vaginal and operative deliveries and were therefore ineligible for inclusion. Only three studies met the inclusion criteria.

Effects of antibiotic prophylaxis on infectious maternal morbidities in vaginal deliveries involving manual placenta removal

All three eligible studies were retrospective in design and conducted in hospitals in Norway (1999), Bulgaria (1983) and Germany (1988) [21–23]. The study population and intervention were consistent across studies. Antibiotics for prophylaxis were introduced for manual placenta removal after placenta retention was diagnosed and women with a prior history of fever were ineligible for inclusion in the study population. All three studies reported on the pre-specified primary outcome, postpartum endometritis. All secondary outcomes (perineal infection and/or any infection, hospital stay duration, sepsis, hemorrhage >1000 ml, hospital readmissions) were not reported on excluding puerperal fever. Only one of the eligible studies reported on this.

Two studies reported on the antibiotic agent used. Von Rechlin et al. [22] used Mebacid/Sulfamerazine (at least over 10 days, 1 g per 1 day and then 0.5 g daily), while Katsulov [21] used 2 g cefamandole (cephalosporin family) for an unstated short duration. In the third study [23], the antibiotic prophylaxis agent was unnamed and the dosage was not given. The authors reported routine use of antibiotics in the center, with the practice maintained throughout the study period.

Both studies loosely defined endometritis as fever greater than 37.5 °C in addition to clinical evidence (not clearly stated in either paper, though probably based on clinician’s judgment) and puerperal fever was defined as a persistent or uninterrupted temperature record of 37.5 °C. The third study was published in English and better defined endometritis as a temperature of 38.0 °C, excessive uterine tenderness or elevated C-reactive protein (±50 mg/l).

Von Rechlin et al. [22] reported a comparable incidence of fever in both the prophylaxis group (89.2 %) and the control group (89.9 %). The other two studies [21, 22] reported only on endometritis as an outcome. Both studies showed no difference between the intervention and control group. Katsulov [21] showed no incidence of endometritis in the prophylaxis group (0 %) compared to control (10.1 %), although this difference was not significant. Tandberg et al. [23] detected a similar incidence of
endometritis in both the prophylaxis (1.6 %) and control (1.9 %) groups.

Follow-up periods ranged from the period of administration of antibiotics (6 days) to the 1st month postpartum, although the period was unstated in Von Rechlin et al. [22].

A meta-analysis similarly showed no difference between prophylaxis and non-prophylaxis groups. For puerperal fever, no difference was observed between groups (odds ratio (OR) 0.93, 95 % confidence interval (CI) 0.38 to 2.27, one study, 302 women).

Similarly, incidence of endometritis was similar in both groups (OR 0.84, 95 % CI 0.38 to 1.85, three studies, 567 women) (Fig. 2). No adverse effects or other secondary outcomes were reported, and no heterogeneity was observed across studies.

Assessment of eligible studies using the Newcastle Ottawa Scale generally showed a low quality of evidence (Table 2), and both outcomes (endometritis and fever) were judged very low in overall quality using the GRADE tool (Table 3).

### Table 2

| Study ID | Representative of exposed cohort | Selection of non-exposed cohort | Ascertainment of exposure | Demonstration that outcome of interest was not present at the start of the study | Comparability of cohorts on the basis of the design or analysis | Follow-up long enough for outcomes to occur | Adequacy of follow-up of cohorts | Total number of stars |
|----------|----------------------------------|---------------------------------|---------------------------|-----------------------------------------------------------------|---------------------------------------------------------------|-----------------------------------------|----------------------------------|---------------------|
| Von Rechlin et al. 1988 [22] | - Truly representative of the average woman (*) | - Drawn from the same community as the exposed cohort (*) | - Secure records (e.g., surgical records) (*) | - Yes (*) | - Study controls for gestational age and/or birth weight (*) | - Yes (*) | - Complete follow-up – all subjects accounted for (*) | 2 |
| Tandberg et al. 1999 [23] | - Somewhat representative of the average woman (*) | - Drawn from a different source | - Structured interview (*) | | - Study controls for any additional factor (*) | | - Subjects lost to follow-up unlikely to introduce bias, or description provided of those lost (*) | 5 |
| Katsulov 1983 [21] | - Selected group of users-No description of the derivation of the cohort | - No description of the derivation of the non-exposed cohort | - Written self-report | | | | | 3 |

**Prophylactic antibiotic regimen for reduction of maternal infectious morbidities during manual placenta removal in vaginal birth**

No eligible studies were found comparing antibiotic classes or regimens for prophylaxis in vaginal deliveries involving manual placenta removal.

**Discussion**

**Main findings**

No significant benefit was observed for the routine use of prophylactic antibiotics on the reduction of maternal febrile morbidities or postpartum endometritis for manual placenta removal. However, the low quality of evidence limits definite conclusions.

Included studies were comparable in terms of study population, intervention and outcome. All three studies individually showed no difference between the control and experimental groups for both fever and endometritis outcomes.

Meta-analysis further showed a similar incidence of reported maternal morbidities (fever and endometritis)
| Quality assessment | Ab prophylaxis versus none | Control | Effect | Quality | Importance |
|--------------------|-----------------------------|---------|--------|---------|------------|
| No. of studies     | Design                      | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | No of patients | Relative (95% CI) | Absolute | |
| Puerperal fever    | 1 observational studies     | very serious a | no serious inconsistency | no serious indirectness | serious b | none | 58/65 (89.2 %) | 213/237 (89.9 %) | OR 0.93 (0.38 to 2.27) | 7 fewer per 1000 (from 127 fewer to 54 more) |
|                    |                             |                     |                    |                    |            |       | 89.9 % | 7 fewer per 1000 (from 127 fewer to 54 more) | VERY LOW |
| Endometritis       | 3 observational studies     | very serious a | no serious inconsistency | no serious indirectness | serious b | none | 8/156 (5.1 %) | 31/411 (7.5 %) | OR 0.84 (0.38 to 1.85) | 11 fewer per 1000 (from 45 fewer to 56 more) |
|                    |                             |                     |                    |                    |            |       | 7.1 % | 11 fewer per 1000 (from 43 fewer to 53 more) | VERY LOW |

Question: Antibiotic prophylaxis versus none for manual placental delivery during vaginal birth
Settings: Hospitals in Germany, Norway and Bulgaria

aStudy contributing data had design limitations based on Newcastle Ottawa Scale rating, non-randomized data collection
bWide confidence interval crossing the line of no effect
between groups. In the random effects analysis, the Von Rechlin study weighted 81% of the pooled weight and appears to favor the use of antibiotic prophylaxis compared to no prophylaxis, though this should be interpreted with caution due to the study's small sample size and low quality of evidence.

**Strengths and limitations**

No evidence exists to date that assesses routine antibiotic prophylaxis for manual placental removal in vaginal birth. This systematic review provides comparable alternative evidence to currently lacking randomized controlled studies and may represent the only systematic evidence available on the subject.

The detailed search strategy, methodology, statistical analyses, absence of statistical heterogeneity and language restrictions, however, lends credence to this review.

Inclusion of studies published in languages other than English ensured the inclusion of important studies on the subject.

This review was limited to evidence from a small number of low quality non-randomized studies and few outcomes due to a clear absence of randomized studies on the subject. Three studies met the inclusion criteria and reported on only two outcomes. Collated data was from observational studies conducted in two developed countries (Germany and Norway) and one developing country (Bulgaria) and may not be globally representative or applicable as policies on antibiotic administration may vary across income settings. The lack of heterogeneity across studies further adds to the validity of the study.

Routine antibiotic prophylaxis is increasingly associated with, antibiotic resistance [24]. Its administration needs to be assessed in the light of emerging resistance and inherent benefits to justify and balance recommendations for routine use; however, eligible studies included in this review failed to report on these points.

In addition, no reports were found on other outcomes in any of the eligible studies or any data comparing the use of different types of prophylactic antibiotics for manual placenta removal in vaginal deliveries. This restricted our evaluation of the effectiveness of routine antibiotic prophylaxis or the definition of an optimal regimen. Studies aiming to determine optimal doses may prove helpful in the future.

**Conclusion**

Available evidence is insufficient to support routine antibiotic prophylaxis for manual placental removal following vaginal birth. This study highlights the need for robust randomized control studies in both low- and high-income settings that will incorporate relevant outcomes such as adverse effects of prophylaxis. Knowledge of this may help inform better assessment of the efficacy of routine antibiotic prophylaxis in obstetric vaginal procedures. The decision to administer antibiotics for manual placenta removal should be selective and based on a balance between the inherent value to the patient and the clinician experience in the light of increasing antibiotic resistance.

Increase in antibiotic resistance in recent years highlights the need for selective prophylaxis to be integrated into best practice and policy against a background of existing area-specific norms and policies.

Successful prevention of maternal infectious morbidities and mortalities resulting from manual removal of the placenta in vaginal births may prove invaluable towards maternal health improvement.

**Abbreviations**

MEDI LNE: Medical Literature Analysis and Retrieval System Online; EMBASE: Excerpta Medica Database; CINAHL: Cumulative Index of Nursing and Allied Health Literature; CI: Confidence Interval; NOS: NewCastle-Ottawa Scale; WHO: World Health Organisation; OR: Odds Ratio; PRISMA: Preferred Reporting Items for Systematic Reviews and Meta-analyses GRADE, Grades of Recommendation, Assessment, Development and Evaluation; MOOSE: Meta-analysis of Observational Studies in Epidemiology.

**Competing interests**

The authors have no conflicts of interests or financial disclosures to declare.

**Authors’ contributions**

Hand-searching, screening, data extraction, meta-analyses, and drafting of the manuscript were conducted by CEC and AJQP. TS assisted with search strategy and conducted the search. EO and OT0 gave guidance on project design, data extraction, meta-analysis and methodological assessments. EO, OT0 and RM assisted in manuscript revision. All authors read and approved the final manuscript.

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