Postal recruitment for genetic studies of preterm birth: A feasibility study [version 1; peer review: 2 approved, 2 not approved]

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Abstract

Background: Preterm birth (PTB) represents the leading cause of neonatal death. Large-scale genetic studies are necessary to determine genetic influences on PTB risk, but prospective cohort studies are expensive and time-consuming. We investigated the feasibility of retrospective recruitment of post-partum women for efficient collection of genetic samples, with self-collected saliva for DNA extraction from themselves and their babies, alongside self-recollection of pregnancy and birth details to phenotype PTB.

Methods: 708 women who had participated in the OPPTIMUM trial (a randomised trial of progesterone pessaries to prevent PTB [ISRCTN14568373]) and consented to further contact were invited to provide self-collected saliva from themselves and their babies. DNA was extracted from Oragene OG-500 (adults) and OG-575 (babies) saliva kits and the yield measured by Qubit. Samples were analysed using a panel of Taqman single nucleotide polymorphism (SNP) assays. A questionnaire designed to meet the minimum data set required for phenotyping PTB was included. Questionnaire responses were transcribed and analysed for concordance with prospective trial data.

Results: Recruitment rate was 162/708 (23%) for self-collected saliva samples and 157/708 (22%) for questionnaire responses. 161 samples from the mother provided DNA with median yield 59.0µg (0.4-148.9µg). 156 samples were successfully genotyped (96.9%). 136 baby samples had a median yield 11.5µg (0.1-102.7µg); two samples failed DNA extraction. 131 baby samples (96.3%) were successfully genotyped. Concordance between self-recalled birth details and prospective birth details ranged from 55 – 99%, median 86%. The highest rates of concordance were found for mode of birth (154/156 [99%]), smoking status (151/157 [96%]) and...
ethnicity (149/156 [96%]).

**Conclusion:** This feasibility study demonstrates that self-collected DNA samples from mothers and babies were sufficient for genetic analysis but yields were variable. Self-recollection of pregnancy and birth details was inadequate for accurately phenotyping PTB, highlighting the need for alternative strategies for investigating genetic links with PTB.

**Keywords**

premature birth, fetal membranes, premature rupture, obstetric labour, premature, surveys and questionnaires, phenotype, DNA

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Introduction
Preterm birth is the leading cause of neonatal morbidity and mortality, resulting in an estimated economic burden to the public sector in England and Wales in excess of £2.9 billion over 18 years\(^1\). Spontaneous preterm birth (PTB) refers to birth less than 37 weeks gestation after the spontaneous onset of contractions\(^2\). In England in 2011/2012 27,509 babies were spontaneous PTBs, of which 11,480 were less than 32 weeks gestation\(^3\). Although our knowledge surrounding PTB and thus our ability to treat and prevent it has been increasing over time, 95% of preterm births are intractable to current therapies\(^4\). Thus, further research into the pathogenesis of PTB is required to decrease this public health problem.

Research has shown that genetic factors contribute to spontaneous PTB. The strongest risk factor for PTB\(^5\), with a recurrence rate after one spontaneous PTB of 15%, which further increases the earlier the gestation, suggesting a maternal genetic component to the risk. Another suggestion of genetic association is the significant ethnic differences in incidence of PTB, with higher rates in women classified as black, African-American and Afro-Caribbean compared with white women, even when environmental confounding factors are taken into consideration\(^6\). A familial predisposition has also been shown, with women who were born preterm being more likely to have preterm babies themselves\(^7\), as well as women being more likely to have PTB if their sisters have had PTB\(^8\). Recent advances in genetic and bioinformatic technologies now provide the potential to understand the complex interaction of genetic and environmental factors. However, studies of genetic associations with pregnancy complications are dependent on very large numbers of good quality DNA samples from well-phenotyped cases, preferably with samples from mother and baby pairs. Studies of genetic associations with conditions such as breast cancer and diabetes have successfully used postal recruitment, with participants donating DNA through provision of saliva samples, which can be returned by post\(^9\). This method could be an efficient way of sample collection from women who have had a PTB and their babies. However, it is crucial to know whether high quality phenotypic information can be provided alongside this approach. An international collaboration of researchers interested in genetic epidemiology studies of pregnancy has highlighted prerequisite phenotypic information essential for performing genetic association studies of preterm birth\(^10\). The group highlights the important differences between spontaneous PTB (spontaneous onset of contractions), spontaneous PTB with preterm premature rupture of membranes (PPROM), and medically indicated PTB (medical indications being fetal compromise, such as small for gestational age, or maternal compromise, for example severe pre-eclampsia)\(^11\). The genetic associations and pathophysiology underlying these three conditions vary greatly\(^12\), hence the necessity for accurate phenotyping in genetic studies in this field specifically. Furthermore, it has not yet been established if maternal recall would be of sufficient quality to support such research. It is also unknown whether postal recruitment and self-collection of samples from mothers, and maternal collection of samples from infants, would be acceptable to participants or yield sufficient quantity and quality of DNA.

There are three aims of this study, which was completed in collaboration with mother and baby pairs from women who took part in the OPPTIMUM trial\(^12\). Firstly, to pilot the method of postal recruitment and sample collection, return and processing. Secondly, to confirm that maternally collected saliva samples, particularly from infants, provide sufficient high quality DNA yield. Thirdly, to assess the agreement of self-recalled pregnancy and birth details in a questionnaire, including essential information for preterm birth genetic association studies, compared with prospectively collected OPPTIMUM trial data.

Methods
Participants
We included UK based participants from the OPPTIMUM trial, a double-blind, randomised, placebo-controlled trial investigating the effect of vaginal progesterone on pregnancy and infant outcomes in women at high risk of spontaneous PTB (https://doi.org/10.1186/ISRCTN14568373)\(^13\). OPPTIMUM recruited from 65 UK National Health Service (NHS) Hospitals and 1 Swedish hospital between February 2009 and April 2013. The inclusion criteria for the OPPTIMUM trial were: high risk for PTB, gestation established by scan at ≤16 weeks to ensure that estimated date of delivery is accurate, signed consent form and aged 16 years or older (no upper age limit). Exclusion criteria for the OPPTIMUM trial were: known significant congenital structural or chromosomal fetal anomaly, known sensitivity, contraindication or intolerance to progesterone, suspected or proven rupture of the fetal membranes at the time of recruitment, multiple pregnancy, prescription or ingestion of medications known to interact with progesterone and women who were prescribed progesterone who took progesterone beyond 18 weeks gestation (See extended data: Appendix 2\(^14\)). For this study, we excluded women who: had withdrawn their consent from the OPPTIMUM trial, those of whom we had no contact details, those whose babies had died subsequent to the 2-year follow-up period, and those who were recruited in Sweden (Figure 1).

Recruitment
Potential participants were sent a letter of invitation to the “OPPTIMUM genetics” study from 20\(^\text{th}\) April 2015 to 23\(^\text{rd}\) July 2015 (see extended data\(^15\)). Women who had a stillbirth, neonatal or infant death were sent an alternate letter of invitation (see extended data\(^16\)). Women were asked to reply by email, phone, text or post to indicate their interest in participating. The study lasted for 12 months after the last recruitment pack was sent out.

Sample collection
Women who responded positively to the invitation letter were sent a recruitment pack (see extended data: Appendix 3\(^17\)). Women who had a live infant were invited to provide a saliva sample from themselves and their baby born whilst participating in the OPPTIMUM study; for those women who had experienced infant loss, an alternative pack was sent with a
collection kit for themselves only. The recruitment packs contained a recruitment letter, patient information leaflet, maternal consent form (2 copies), child assent form (if child over 4 years old), instructions for saliva sample collection for mother and maternal saliva sample collection kit (Oragene OG-500, DNA Genotek), a clinical data questionnaire and a postal return kit (postage paid; see extended data 13). Where appropriate, instructions for saliva sample collection for baby, and an infant saliva sample collection kit (Oragene OG-575, DNA Genotek) was included. If the participant’s baby had died, they were invited to provide a saliva sample from themselves, and an alternative recruitment pack was sent (see extended data: Appendix 313). Women who had not returned the recruitment pack after 6–8 weeks were sent a single reminder letter.

**Figure 1. Study flow diagram of recruitment.**

**Questionnaire design and application**

The questionnaire was designed by the OPPTIMUM trial team to meet the minimum data required for a study of PTB14 and was piloted in 20 postpartum women. It included questions on gestation at birth, maternal age at birth, birth method, labour onset, membrane rupture, maternal smoking, non-prescription drug use and alcohol intake, as well as the number of previous pregnancies and maternal ethnic origin (see extended data: Appendix 113). Participants were asked to answer the questions in relation to their pregnancy during the OPPTIMUM trial.

**Receipt, processing and storage of samples**

The Edinburgh Clinical Research Facility Genetics Core received and processed the samples. Samples were identified by the
OPPTIMUM trial number, with a suffix for mother and baby and labelled with details readable by barcode scanner.

DNA extraction and validation
DNA was extracted from Oragene OG-500 (adults) and OG-575 (babies) saliva kits using Oragene prePT (PT-L2P-5) extraction kit (supplied by DNA Genotek). DNA yield was measured by Qubit (ThermoFisher). Samples were genotyped on a panel of Taqman single nucleotide polymorphisms (SNPs) using the QuantStudio12K Flex and analysed using QuantStudio v1.2.2 software. Samples from mothers were genotyped using autosomal SNPs rs6427699, rs4751955, rs11083515, rs7588807, rs10938367 & rs10869955. Samples from babies were run on the same six autosomal SNPs and an additional three SNPs from the Y-chromosome to determine sex (rs2032598, rs768983 & rs3913290). An aliquot of DNA was normalised in plates for future analysis. DNA samples were transferred for storage and used as part of the Edinburgh Reproductive Tissues Biobank (REC reference 09/S0704/3).

Data analysis
For this study, data was transcribed from the questionnaires and appropriate information was obtained from the trial database. Patient identifiable information was removed and trial data was correlated with the corresponding questionnaire through randomised OPPTIMUM trial numbers. In keeping with the Caldicott principles14, access to trial data was only granted to members of the research team, and stored on a password protected database on a secure server (University of Edinburgh). The concordance between self-recalled birth details and prospective trial birth details was then analysed for gestation at birth, maternal age at birth, mode of birth, onset of labour, smoking, non-prescription drug use, alcohol use and number of previous pregnancies. The concordance for PPROM was analysed in women who had a PTB. We pre-specified that a participation rate of ≥ 50% would be acceptable and concordance of ≥ 95% between self-recalled birth details and prospective trial birth details would be sufficient for phenotyping PTB. These values were chosen by the research team as previous studies of self-collected DNA samples have had participation rates above 50%15,16,17 and accurate recall using patient questionnaires would be necessary for any future large scale studies using this design.

Statistical analysis
All data was analysed using IBM Statistical Package for the Social Sciences (SPSS) Version 22. Normally distributed data was analysed using a t-test. The Fishers exact test was used for proportional data. A p-value of <0.05 was considered significant.

Ethical opinion
This study was awarded a favourable ethical opinion by the regional ethics committee (REC reference 14/SS/0086).

Results
Recruitment rate and participant demographics
In total, 708 women were contacted. From these, 157 questionnaires were returned, a participation rate of 22%. Overall, 299 DNA sample kits were received (137 mother and baby paired samples, 24 mother only samples, 1 baby only sample) - a participation rate of 162/708 (23%) (see Figure 1). Seven participants returned a DNA sample kit without a questionnaire and 2 participants returned the questionnaire without a DNA sample. One participant was recruited twice to the OPPTIMUM trial and so returned a questionnaire for each pregnancy, a mother and baby paired sample from her first child and a baby only sample from her second child. See underlying data for all data collected13.

The demographics of the questionnaire respondents are shown in Table 1. To determine if these 157 women were representative of the OPPTIMUM trial population the demographics of the participants in this pilot study were compared to that of the entire cohort of OPPTIMUM trial participants. This highlighted that the pilot study participants were significantly older, more educated, had a higher proportion of white participants, and a lower proportion of black participants compared with the entire OPPTIMUM cohort (Table 1).

DNA extraction and quality assessment
DNA was extracted from 299 saliva samples. All 161 samples from the mother provided DNA with median yield 59.0µg (0.4–148.9µg). Two baby samples failed and provided no DNA. The remaining 136 baby samples had a median yield 11.5µg (0.1–102.7µg) (Table 2). Samples were genotyped using a panel of six autosomal Taqman SNPs and were deemed suitable for genomic analysis if they successfully genotyped on five or more SNPs. 156 of the 161 samples (96.9%) from the mother successfully genotyped (one sample failed on four SNPs, three samples failed on five SNPs and one sample failed on all six SNPs). 131 of the 136 baby samples (96.3%) successfully genotyped (three samples failed on five SNPs and two samples failed on all six SNPs) (Table 2).

The baby samples were additionally genotyped on three Y-chromosome SNPs. Samples that successfully called on five or six autosomal SNPs were checked for sex, with samples having a no-call on all three Y-chromosomes SNPs determined as female and samples having a call on all three Y-chromosome SNPs determined as male. Samples having a positive call on only one or two Y-chromosome SNPs were assigned to ‘unknown’, of which 6 of 151 samples were assigned. Additionally, 2 samples did not have matched trial data to verify sex. In total, 121 of 124 samples (97.6%) correctly identified the sex of the baby; 3 samples which were assumed female from genotyping were male (Table 2).

Concordance of self-recalled birth details and prospective trial birth details
Concordance of self-recalled birth details and prospective trial birth details for each variable are displayed in Table 3 and Figure 2. Concordance ranged from 55% for PPROM to 99% for mode of birth. Median concordance was 86%. Overall, concordance of self-recalled birth details and prospective trial birth details was only greater than 95% in 3 out of 10 key fields for phenotyping PTB (mode of birth, smoking status and ethnicity).
Table 2. Results of DNA analysis and genotyping from mother and baby self-collected saliva samples.

<table>
<thead>
<tr>
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<th>Mother samples (n=161)</th>
<th>Baby samples (n=138)</th>
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<tr>
<td>Number of samples suitable for analysis</td>
<td>161 (100%)</td>
<td>136 (98.6%)</td>
</tr>
<tr>
<td>Median DNA yield in μg (range)</td>
<td>59.0 (0.4-148.9)</td>
<td>11.5 (0.1-102.7)</td>
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<tr>
<td>Number of samples successfully genotyped using 6 autosomes</td>
<td>156/161 (96.9%)</td>
<td>131/136 (96.3%)</td>
</tr>
<tr>
<td>Number of samples correctly identifying sex</td>
<td>121/124 (97.6%)</td>
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For the variable ‘gestation at birth’, concordance between self-recalled birth details and prospective trial birth details was improved in earlier gestations: 15/18 (83%) concordance for birth under 34 weeks, 27/34 (79%) for birth between 34–36 weeks inclusive and 76/101 (75%) for birth at 37 weeks or later. 144/153 (94%) of respondents self-recalled a gestation of birth within 1 week of the data in the prospective trial and 149/156 (96%) self-recalled their age at birth within 1 year of the prospective trial data.

Preterm birth

In prospectively collected trial data 52/157 (33%) respondents had a PTB. Of these, 15/52 (29%) women had a medically indicated preterm birth (i.e. induced labour, or elective caesarean section due to suspected fetal compromise, maternal condition or previous caesarean section). Onset of labour was spontaneous in the remaining 37/52 (71%) of women. In the prospectively collected trial data 11/37 (30%) of spontaneous PTBs were preceded by PPROM. There was concordance in the presence of PPROM between prospective trial birth details and self-recalled birth details in only 6/11 (55%) cases (Figure 2).

Smoking status, non-prescription drug use, alcohol consumption

There was no statistical difference found between the number of smokers reported in prospective trial data and self-recalled data (p=0.838). The difference in non-prescription drug use...
Table 3. Concordance of prospective trial birth details with self-recalled pregnancy and birth details.

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<th>Table 3. Concordance of prospective trial birth details with self-recalled pregnancy and birth details.</th>
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<tr>
<td><strong>Mode of delivery</strong></td>
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<tr>
<td><strong>Smoking status</strong></td>
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<tr>
<td><strong>Ethnicity</strong></td>
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<tr>
<td><strong>Labour onset</strong></td>
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<tr>
<td><strong>Drug use</strong></td>
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<tr>
<td><strong>Alcohol use</strong></td>
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<td><strong>Maternal age at birth</strong></td>
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<td><strong>Gestation at birth</strong></td>
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<tr>
<td><strong>Number of previous pregnancies</strong></td>
</tr>
<tr>
<td><strong>Preterm premature rupture of membranes</strong></td>
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</table>

Figure 2. Concordance of prospective trial birth details with self-recalled pregnancy and birth details for each of the questions in the participant questionnaire.
between prospective trial data and self-recalled data was statistically significant (p=0.0001); 18/157 (11%) of women recorded as no drug use in prospective trial data classified themselves as having used non-prescription drugs during their pregnancy. A subset of these women (10/18, 56%) included the names of the drugs used; these included paracetamol, aspirin, folic acid, codeine and “indigestion medication”. The difference between alcohol consumption in prospective trial data and self-recalled data was statistically significant (p=0.0005); 23/148 (16%) women recorded as non-drinkers in prospective trial data self-recalled use of alcohol during their pregnancy.

Number of previous pregnancies
With regards to number of previous pregnancies, there was a discrepancy in 53/157 (34%) cases. In 30/53 (57%) of these, a higher number of previous pregnancies was recorded in prospective trial data than in self-recalled data.

Discussion
The primary aim of this study was to pilot postal donation of self-collected DNA samples from postpartum women and their babies. The recruitment rate of 23% was below our pre-specified response rate of ≥ 50% and compares poorly with other studies investigating self-collection of DNA which have had participation rates greater than 50%.[5,12]. Only one reminder was sent - it is possible that further reminders might have increased the participation rate. We recruited participants from a cohort of women who had previously taken part in the OPPTIMUM trial. Although this limited our study population, the main advantages were that consent had already been obtained and that high quality prospectively collected data on demographics and birth outcomes was available for validation.

The Preterm Birth Genome Project investigated multiple methods of DNA collection from mother and baby pairs (whole blood, blood spot, buccal, saliva) from four countries and found that samples were not affected by transportation methods and that salivary samples provided an adequate yield of DNA, superior to buccal swabs.[8] Similar studies of self-collected saliva samples report DNA yields of > 70%.[9,20] Our study shows that postal self-collected saliva samples from mothers and babies were of sufficient quality for genetic analysis but yields were variable.

We also aimed to determine the concordance between self-recalled birth details and prospective trial birth details with a view to determine if postal questionnaires are a valid method of phenotyping PTB. We demonstrate that concordance was above the pre-specified 95% in only three out of ten questions. It is very unlikely that the prospectively collected OPPTIMUM trial data was erroneous. The study was carried out to rigorous clinical trial standards with a pre-defined data dictionary and training of all research staff contributing to data collection. There was regular trial data monitoring from the sponsor ensuring data quality and consistency with checking of source data.

Labour onset and membrane rupture are important features which differentiate spontaneous PTB from PTB following PPROM and medically indicated PTB. This is crucial, as the underlying genetic basis for each phenotype is potentially different. In this dataset, concordance for labour onset was reasonably good at 92%. We assessed concordance for the occurrence of PPROM in women who had a PTB, which was low (despite wide confidence intervals), suggesting that PPROM is not accurately recalled. However, we recognise that this is a very small subset of women and a larger number would be required to accurately assess this finding.

We have demonstrated poor concordance between self-recalled birth details and prospective trial birth details with regards to age at birth, gestation at birth and number of previous pregnancies. Of note, the index birth was up to five years ago. Despite piloting and validating questionnaires, the accurate recall of time dependent features are still susceptible to recall bias and so this method may not be reliable for gathering critical information. Answers to such questions may also be dependent on the level of health or mathematic literacy in the study population, though the mean period in full-time education for this cohort was 14.4 years. In addition, the question asking about previous pregnancies fell on a page break which split the question (see extended data: Appendix 1[13]). This led to some women incorrectly completing the question, confounding our results. This emphasises the importance of piloting a questionnaire in its completed, printed form.

Compared to the high level of concordance for smoking status, the concordance between prospective trial data and self-recalled data was much lower with regards to non-prescription drug use and alcohol intake: a much higher level of alcohol consumption was reported in self-recalled data than in prospective trial data. Interestingly, the discrepancies between the reporting of smoking, alcohol and non-prescription drug use are in both directions. This highlights differences in public perception and willingness to disclose such information whilst in a research trial or in an anonymised questionnaire. The inconsistencies in reporting of non-prescription drug use were mainly due to several women self-recalling use of ‘over the counter’ medications such as paracetamol. This is in contrast to the OPPTIMUM trial definition of non-prescription drug use which included: heroin, cocaine or abuse of prescribed drugs such as benzodiazepines. This finding is most likely due to a misinterpretation of the question and further highlights the importance of accurate wording in similar questionnaire studies.

In conclusion, this feasibility study shows that women can successfully collect DNA from themselves and their babies, but overall yields were variable. Yield from babies was lower than mothers due to the lower amount of saliva collected with the Oragene OG-575. Taqman genotyping showed the samples were suitable for variant calling and checking sex. The low yields, particularly for the baby samples, would make some methods of genomic analysis challenging, such as exome sequencing, but with the development of low-input kits even these methods
may be suitable. Consideration should be given to the genetic analysis to be performed when deciding on a collection method.

We found postal post-partum participation rates are low and there were significant discrepancies between self-recall of pregnancy and birth details and prospective birth details as recorded in the OPPTIMUM trial dataset. We conclude that information gathered from postal questionnaires is insufficient to accurately phenotype PTB and clinical data collection from medical records needs to be an integral part of any future study design into the genetics of preterm birth.

Data availability

Underlying data

This project contains the following underlying data:
- Figshare1. DNA data.csv (Dataset showing total DNA yield for all self-collected samples)
- Figshare2. OPPTIMUM trial data.csv (Prospective trial data)
- Figshare3. Questionnaire data.csv (Data from participant questionnaires)
- Figshare4. Comparison results of y markers_281019_BABY.csv (Results of determining sex of baby from saliva samples)
- Figshare5. Taqman Genotyping Report_291019.pdf (Report detailing results of Taqman genotyping)
- Genotyping results for 6 x SNPs using mother and baby samples (individual files for Taqman genotyping results for mother and baby for each SNP: rs6427699, rs4751955, rs11083515, rs7588807, rs10938367 & rs10869955)

Extended data

This project contains the following extended data:
- Appendix 1. Participant questionnaire (example of participant questionnaire)
- Appendix 2. OPPTIMUM trial inclusion and exclusion criteria (list of inclusion and exclusion criteria for recruitment to the OPPTIMUM trial)
- Appendix 3. List of recruitment pack contents (list of recruitment pack contents for women who had a live birth, and for women who had a stillbirth, neonatal or infant death)
- OPPTIMUM Genetics invitation letters (invitation letters to participate in OPPTIMUM Genetics study, one for women who had a live birth, one for women who had a stillbirth, neonatal or infant death)
- Recruitment pack letters (letters included in recruitment pack, one for women who had a live birth, one for women who had a stillbirth, neonatal or infant death)
- OPPTIMUM genetics patient information leaflets (patient information leaflets, one for women who had a live birth, one for women who had a stillbirth, neonatal or infant death)
- OPPTIMUM genetics consent forms (participant consent form, one for women who had a live birth including consent for her baby or child to participate, one for women who had a stillbirth, neonatal or infant death)
- OPPTIMUM genetics child assent form (for children over 5 years of age to assent to participation in study)
- Saliva sample collection instructions (instructions on how to collect saliva sample from participant and from baby or child using Oragene saliva kits)

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Acknowledgements

The authors would particularly like to thank Amy Witherspoon, Tariq Derdeb, Lorraine Adamson and Heather Macpherson who were involved with data and sample collection, William Hawkins for providing the Taqman genotyping, and all of the OPPTIMUM trial participants and research team.

References

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PubMed Abstract | Publisher Full Text

15. Chairman’s foreword: Information: To share or not to share? 2013. 
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Version 1

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Ge Zhang
Division of Human Genetics, The Center for Prevention of Preterm Birth, Perinatal Institute, Cincinnati Children’s Hospital Medical Center, Cincinnati, OH, USA

This pilot study evaluated the feasibility of retrospective collection of DNA samples and self-recalled phenotype data for genetic studies of preterm birth. The study demonstrated that the DNA samples were sufficient for genetic analysis but the self-recalled phenotype data was inaccurate. The reviewer has major concern with using concordance in evaluating the consistency of phenotype data.

1. Introduction: ref 11 demonstrated that successful genetic studies of gestational duration and preterm birth require large sample sizes. It did not study the genetic or pathophysiological differences among sPTB, PPROM and indicated PTB.

2. Methods (Data analysis): The calculation of concordance should be explicitly explained.

3. Methods (Data analysis): In addition to concordance, the authors should check correlations (especially for quantitative variables and binary variables) between the self-recalled data and data collected from the clinical trial.

4. Results (first paragraph): The authors may provide Venn diagrams to show the overlapping of DNA samples between mothers and children [24 (137) 1] and overlapping between questionnaires and DNA samples [2 (155) 7].

5. Results (second paragraph and Table 1): It should be clarified that the numbers [n(%)] were calculated based on trial data even for the pilot study.

6. Results (DNA quality assessment): The author should present Mendelian consistency between mothers and their child.

7. Results (Table 3): concordance of PPROM was calculated as 6/11 (55%), which might be incorrect (please also see my comment 2).
8. Results (Please also see my comments 2 and 3): It seems to the reviewer that concordance is not an adequate measure to evaluate the consistency between self-recalled data and the data collected during the trial. For example, only 1 mother in the pilot study was recorded having non-prescription drugs; but 18 recorded as no drug use recalled themselves as having used non-prescription drugs. This information is totally inconsistent and misleading but still the concordance is 88% = (155-(18+1))/155. The same is true for alcohol consumption. As suggested by my comment 3, the authors should check correlations or present Venn diagrams for every categorical variable to show the complete picture of relationships between the two data sets.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Statistical genetics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.

Reviewer Report 15 April 2020

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Kelly-Ann Eastwood
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This is an interesting study investigating feasibility of self-collected postnatal maternal and neonatal saliva for DNA extraction. The paper is well written and of interest to clinicians and researchers.

The authors have clearly discussed limitations of the study, including recruitment rate (23%) and poor concordance between self-recalled birth details and prospective trial data.

I wonder if recruitment rate may have in fact been bolstered by previous participation in the OPTIMMUM study. This could have implications in a real world clinical setting. Regarding collection of saliva samples; median DNA yield demonstrated a large range for both mother and neonatal samples. Going forward, how could integrity of sampling technique be assured?

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate? I cannot comment. A qualified statistician is required.

Are all the source data underlying the results available to ensure full reproducibility? Yes

Are the conclusions drawn adequately supported by the results? Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Academic Clinical Lecturer in O&G, sub-specialty trainee in maternal and fetal medicine. Research interests include pre-eclampsia, pre-conception care and maternal obesity.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
This study examines the feasibility of postnatal recruitment for analysis of genetic factors associated with preterm birth. It is methodologically appropriately designed and the conclusions drawn by the authors seem appropriate. This study highlights the difficulty involved in reliance upon recruitment into studies remotely and the problems with self reporting of outcomes which often don't tally with correct medical diagnosis.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
I cannot comment. A qualified statistician is required.

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Maternal and Fetal medicine, preterm labour, multiple pregnancy

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Tessa Homfray
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- I am uncertain exactly when these women were recruited. The OPPTIMUM study was 2013 and therefore this was sometime later that this was attempted which would I suspect lead to the poorer recall than if this had been attempted later.
The amount of baby saliva DNA would be too small to be of much use I suspect.

I didn't see any evidence confirming maternity of the fetal samples which would indicate if they had been collected correctly.

This is a proof of principle study which suggests this is not a good method of obtaining genetic material. On this basis I think the article should be much shorter.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Consultant Medical Genetics with Particular interest in fetal medicine.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.