Use of metabolomics for predicting spontaneous preterm birth in asymptomatic pregnant women: protocol for a systematic review and meta-analysis

Renato T Souza,1 Rafael Bessa Galvão,1 Debora Farias Batista Leite, 2,3 Renato Passini Jr,4 Philip Baker,5 Jose Guilherme Cecatti 1,6

ABSTRACT

Introduction Preterm birth (PTB) is the leading cause of neonatal mortality and short- and long-term morbidity. The aetiology and pathophysiology of spontaneous PTB (sPTB) are still unclear, which makes the identification of reliable and accurate predictor markers more difficult, particularly for unscreened or asymptomatic women. Metabolomics biomarkers have been demonstrated to be potentially accurate biomarkers for many disorders with complex mechanisms such as PTB. Therefore, we aim to perform a systematic review of metabolomics markers associated with sPTB. Our research question is ‘What is the performance of metabolomics for predicting spontaneous preterm birth in asymptomatic pregnant women?’

Methods and analysis We will focus on studies assessing metabolomics techniques for predicting sPTB in asymptomatic pregnant women. We will conduct a comprehensive systematic review of the literature from the last 10 years. Only observational cohort and case-control studies will be included. Our search strategy will be carried out by two independent reviewers, who will scan title and abstract before carrying out a full review of the article. The scientific databases to be explored include PubMed, MedLine, ScieLo, EMBASE, LILACS, Web of Science, Scopus and others.

Ethics and dissemination This systematic review protocol does not require ethical approval. We intend to disseminate our findings in scientific peer-reviewed journal, the Preterm SAMBA study open access website, specialists’ conferences and to our funding agencies.

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INTRODUCTION

Spontaneous preterm birth (sPTB) is the leading cause of perinatal mortality and short- and long-term morbidity.1 It is defined as birth that occurs before 37 weeks gestation due to spontaneous onset of labour or preterm premature rupture of membranes (pPROM).3 4 Several pathways and mechanisms linked with PTB have been proposed including, neuroendocrine, vascular, immune-inflammatory and behavioural processes.3 More specifically, several markers associated with uterine distension/contraction, decidual inflammation/infection and activation of hypothalamic-pituitary-adrenal axis had been studied in the past decades.5 6 However, no single marker or combination of markers has been found to be accurate enough for predicting sPTB.7–10 History of previous PTB, cervical length at second trimester and cervico-vaginal fetal fibronectin (fFN) are the most promising clinical tests for predicting spontaneous preterm, but they seem not to be clinically useful for asymptomatic women. Sensitivity of short cervical length (<25mm) and high cervico-vaginal fFN (>50ng/mL) are around 33%–36% and 46%, respectively.11–13

PTB is a complex and multifactorial syndrome that possibly has a long pre-clinical
phases, maternal and fetal interactions, genetic and environmental influences, and adaptive mechanisms.14 15 These challenging aspects, and the presence of still unknown underlying mechanisms, are the main limitations for the identification of an accurate predictor for sPTB.16–18 None of the predictors used in clinical practice, such as previous history of PTB, infection (vaginal and urinary contaminants), FN and transvaginal ultrasonography cervical length demonstrated exceptional accuracy for predicting sPTB.7 An exploration of innovative approaches is urgently required.

Metabolomics is the study of metabolites, through identification and quantification of low-weight molecular particles, ie, tens to hundreds thousands of intermediate products and substrates of systems biology chemical reactions.19 20 This novel approach has been applied for identifying biomarkers and underlying biochemical pathways associated with complex obstetrical syndromes as pre-eclampsia, fetal growth restriction, gestational diabetes and PTB. In contrast to other ‘Omics Sciences’ techniques, metabolomics is more closely associated with the phenotype of the disease and might thus identify a more robust and reliable set of predictors.21 Importantly, implementing an adequate Omics experimental design is crucial for metabolomics studies. Using different baseline population (asymptomatic vs symptomatic or low vs high-risk women for developing sPTB), study designs (prospective cohorts, case–control or cross sectional studies), sources of samples (amniotic fluid, vaginal fluid, blood, urine, hair, etc) and the timing of sample collection each have significant effects on study findings and the consequent interpretation and contribution to the current gap of knowledge.19

Different reviews collating scientific knowledge regarding PTB biomarkers/predictors has been conducted. Different methodology approaches has been applied so far, including narrative, systematic and umbrella reviews, a more comprehensive review that includes not only original studies but also other reviews.7 22–24 At the best of our knowledge, there is no systematic review on metabolomics markers. Therefore, we aim to conduct a systematic review of original studies investigating the use of metabolomics biomarkers for predicting sPTB in asymptomatic pregnant women. This protocol describes the methods that will be applied in our systematic review.

METHODS AND ANALYSIS

The current systematic review proposal will be conducted, written and published following the Preferred Reporting Items for Systematics Reviews and Meta-Analyses (PRISMA-P) recommendations.25

Review question
What is the performance of metabolomics for predicting sPTB in asymptomatic pregnant women?

Eligibility criteria
Original cohort or case–control studies involving asymptomatic pregnant women at the moment of sample collection (exposure) and with samples analysed using metabolomics techniques. Studies will be excluded if (1) they are cross-sectional studies, clinical trials, editorials, letter to editors, case reports, expert opinions, commentaries or any type of review; (2) they describe only experimental studies with animals; or (3) they are duplicated data (eg, data published in conferences proceedings and, then, published again in scientific journals). In this case, only the most complete publication will be considered, after comparing and confirming that the same technique and metabolites were explored. Studies published from 2008 to 2018 will be considered, and there will be no language restriction. Before submitting this systematic review for publication, we will rerun the search strategy to identify new studies that have been published after performing the first round of search.

Participants
The current review is interested in evaluating the performance of metabolomics biomarkers for sPTB in asymptomatic pregnant women, which may contribute to clinical practice, potentially providing information regarding onset of preterm labour. Nevertheless, we aim to identify studies addressing only early predictors collected from women who are in an early preclinical stage, which might contribute to a wider window of opportunity for interventions and also to develop a widely reproducible screening test. Asymptomatic pregnant women should not have regular uterine tightening/contractions or signs of rupture of membranes (ie, watery discharge). In addition, the study should preferably have a standardised definition of sPTB, the outcome of interest.

Information sources
The search will be held in the following databases: PubMed, EMBASE, Scopus, CINAHL, and Web of Science, BVS/BIREME, which includes the Latin American and Caribbean Health Sciences Literature (LILACS), Medline and the Scientific Electronic Library Online (Scielo). In addition, secondary sources of original studies will be explored such as Google Scholar, hand-held searching of the reference list of eligible studies, conference proceedings, and contact with authors when necessary.

Search strategy
The following terms will be used in our search strategy for the different scientific databases: (preterm birth, premature birth, premature infant, premature labour, extremely premature infant, premature obstetric labour, spontaneous preterm birth, extreme preterm birth, late preterm birth, moderate preterm birth, preterm premature rupture of membranes, preterm delivery, PROM, sPTB, preterm PROM, pPROM, p-PROM) AND (metabolomic*, metabonomic*, metabolit*, lipidomic*, H NMR, proton NMR, proton nuclear magnetic resonance, liquid chromatogra*, gas chromatogra*, UPLC, ultra-performance liquid chromatograph*, ultra-performance liquid chromatograph*, HPLC, high performance liquid
Data management

We will export search results to a reference manager (Mendeley). Then, the following information will be collected from each study using an appropriate form, which will be entered in an Excel spreadsheet: author’s name, year of publication, country, study design, number of participants with and without sPTB, type of metabolomics analysis technique (liquid or gas chromatography, nuclear resonance), laboratory methods for metabolites data acquiring (targeted or untargeted techniques, etc.), subtype of PTB (spontaneous preterm labour or pPROM), number of fetuses (singleton vs multiple), gestational age when samples were collected, source of samples (type/site of tissue), low- or high-risk for PTB (authors criteria used to define the population will be collected) and method applied to estimate gestational age. If possible, additional variables related to sPTB categories (delivery before 28 weeks and before 34 weeks) will be recorded for secondary analyses. Original authors will be contacted to clarify data, when needed. Finally, we will check the biochemical class of identified metabolites in Human Metabolome Database (HMDB, version 4.0) to explore and synthetize whether there are common biological pathways associated with sPTB. 

Selection process

Two independent reviewers (RTS and RBFG) will be responsible for screening and selecting studies initially according to title or abstract. Both researchers will read the full text of non-excluded studies to discriminate eligibility. A third reviewer (DFBL) will consider any disagreement; additional reviewers (RPJ, PNB and JGC) will be responsible for supervising all steps and approving data extraction.

Data collection process

We will extract search results to a reference manager where all studies will be stored. Then, included studies will be placed in a new folder. Finally, we will manually extract data of interest from these included studies to an Excel file. Each reviewer will have their own reference manager account, file and folder and discrepant results will be discussed together with the third reviewer.

Outcomes and prioritisation

The primary outcome is sPTB, defined as any birth occurred before 37 weeks of gestation due to spontaneous onset of labour or pPROM. Secondary outcomes are:
1. sPTB before 28 weeks;
2. sPTB before 32 weeks;
3. sPTB before 34 weeks.

The capacity to predict different degrees of sPTB (categories of gestational age) is important as the extreme (<28 weeks), moderate (<32 weeks) and non-late preterm (<34 weeks) newborns have different adverse outcomes compared with non-extreme (≥28 weeks); non-moderate (≥32 weeks) or late (≥34 weeks) preterm newborns.

Ideally, the method of gestational age estimation should be clearly reported. For instance, it can be reported as estimated by last menstrual period (LMP) and confirmed by an early ultrasound or only by an early ultrasound when LMP is unknown/uncertain.

Index test

Metabolomics techniques to predict sPTB is the diagnostic test of interest. Metabolomics is a technique to identify and quantify metabolites from biological samples using different type of platforms/equipment. The most common platforms include gas, liquid chromatography or ultra-performance liquid chromatography coupled to a mass spectrometer or a proton nuclear magnetic resonance. The performance of the different thresholds of each metabolite will be compared and summarised through hierarchical summary receiver operator characteristic curve (meta-analysis) according to the subgroups described above. Considering that the raw data is not available in the majority of the diagnostic test accuracy studies and that metabolites levels are usually reported as continuous variables, we intend to use a meta-analysis model based on receiver operating characteristic (ROC) curves. Briefly, a two-parameter model, based on the estimation of α and β parameters (using standard errors or maximum likelihood), will be applied as reported by Kester and Buntinx. Therefore, pooled ROC curves can be converted to a estimated ROC curve with 95% CI. This method can also be applied in categorical-ordinal variables tests.

Risk of bias in individual studies

We will apply the Quality Assessment of Diagnostic Accuracy Studies (QUADAS-2) tool to assess the risk of bias and applicability of primary diagnostic accuracy studies. Each study will be classified as ‘low’, ‘high’ or ‘unclear’ regarding risk of bias for each of the four domains of QUADAS tool: Patient Selection, Index Test (metabolomics), Reference Standard (occurrence of PTB) and Flow and Timing of participant’s inclusion and follow-up. For example, studies will be labelled as ‘low’ risk of bias for Reference Standard when definition of sPTB and gestational age estimation are clear; ‘high’ risk of bias would be considered when the moment of sample collection is not well described.

Data synthesis

We will report details of identification, screening, eligibility and included studies using a flow diagram, according to PRISMA recommendations. Data from included studies
will be synthesised into tables according to the variables of interest. If possible, we will present data meta-analysis according to study design, metabolomics technique and type of samples analysed. We intend to perform subgroup analysis according to:

- different metabolomics methods applied: gas or liquid chromatography coupled with mass spectrometry or proton nuclear magnetic resonance;
- singleton and multiple pregnancies;
- low-risk and high-risk women for developing PTB;
- subtype of PTB: sPTB exclusively due to spontaneous onset of labour with intact membranes or sPTB due to PROM;
- gestational age interval when samples were collected: first trimester, second trimester and third trimester.

Heterogeneity will be assessed by Cochran’s Q, Hotelling’s T-squared ($t^2$) and $I^2$ tests. Funnel plots and sensitivity and cumulative analyses will be applied for detection of temporal trends and publication bias.

**Potential anticipated limitations to this review**

First, although we have not considered any language restriction, we consider that there might be a limitation in studies published entirely in non-English language. However, in the last decade, more than 95% of scientific biomedical literature has been published in English, then we consider this a minor selection bias. Second, we intend to stratify the groups according to population risk. However, the characterisation of low- or high-risk for sPTB is controversial and lacks standardisation, which might limit data comparison and subgroup analysis. Finally, categorisation of sPTB into spontaneous onset of labour or pPROM is another topic of potential limitation—the recognition of the main initial mechanism for preterm delivery might not always be possible. Even when specified, it might provoke uncertainty and could limit further considerations regarding preterm phenotypes. In addition, another limitation is that individual patient data will not be collected.

**Patient and public involvement**

Patients will not be directly involved in the study and no experience or direct impact from their perspective can be discussed.

**ETHICS AND DISSEMINATION**

We intend to disseminate our findings in scientific peer-reviewed journal, general free access website of Preterm Screening and Metabolomics in Brazil and Auckland (Preterm SAMBA) study, specialists’ conferences, and to our funding agencies.

**DISCUSSION**

This systematic review will comprise current knowledge related with metabolomics in the context of PTB prediction. Metabolomics science, a resourceful innovative field that allows better understanding on pathophysiology of complex syndromes, may address the main compounds associated with the spontaneous preterm delivery and, therefore, motivate further researchers to validate early measurable predictors of PTB.

Metabolomics performance for predicting sPTB remains unclear and standardised and high-quality studies are needed to clarify the clinical application of metabolites for predicting sPTB. Nevertheless, metabolomics discovery studies commonly requires further validation studies; reproducible methodology is crucial. This systematic review protocol will collate the main potential early biomarkers, subgroup analysis and standardised definition for sPTB to better understand metabolomics performance in predicting sPTB and also to show its heterogeneity in terms of methodology (samples used, metabolomics technique, definition of sPTB phenotype, etc.). High performing predictors of PTB will help combat this leading cause of neonatal mortality and morbidity.

**Author affiliations**

1 Obsetetrics and Gynecology, Universidade Estadual de Campinas, Campinas, Brazil
2 Department of Tocogynecology, Campinas’ State University, Campinas, Brazil
3 Department of Maternal and Infant Health, Universidade Federal de Pernambuco, Recife, Brazil
4 Universidade Estadual de Campinas Faculdade de Ciencias Medicas, Campinas, Brazil
5 University of Leicester, College of Medicine, Leicester, UK
6 Obstetrics and Gynecology, University of Campinas, Campinas, Brazil

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**Contributors**

RTS and RBG conducted the systematic review as independent first reviewers. JGC, RP and DFBL made the conflicting decisions regarding papers selections. PB, RP and JGC participated in the systematic review conception, methodology and framework, together with all the others co-authors.

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**Competing interests**

All authors are carrying original research on metabolomics and presenting conferences about this topic, including spontaneous preterm birth, pre-eclampsia, gestational diabetes mellitus and fetal growth restriction. Philip N Baker is principal investigator of Metabolomics Diagnostics Ltd, a company dedicated to develop innovative screening tests using metabolomics technology.

**Patient consent for publication** Not required.

**Ethics approval**

This systematic review does not require ethical approval from the Research Council or Ethics board.

**Provenance and peer review**

Not commissioned; externally peer reviewed.

**Open access**

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REFERENCES
1. Lawn JE, Cousens S, Zupan J. Lancet Neonatal Survival Steering Team. 4 million neonatal deaths: when? Where? Why? Lancet 2005;365:891–900.
2. Howson CP, Kinney MV, McDougall L, et al. Born too soon: preterm birth matters. Reprod Health 2013;10(Suppl 1):S1.
3. Blencowe H, Cousens S, Chou D, et al. Born too soon: the global epidemiology of 15 million preterm births. Reprod Health 2013;10:S2.
4. Ananth CV, Vintzileos AM. Epidemiology of preterm birth and its clinical subtypes. J Matern Fetal Neonatal Med 2006;19:773–82.
5. Behrman RB. Preterm Birth: Causes, Consequences, and Prevention. Washington, D.C: National Academies Press, 2007.
6. Manuck TA, Esplin MS, Biggio J, et al. The phenotype of spontaneous preterm birth: a clinical application tool. Am J Obstet Gynecol 2015;212:487.e1–487.e11.
7. Honest H, Hyde CJ, Khan KS. Prediction of spontaneous preterm birth: no good test for predicting a spontaneous preterm birth. Curr Opin Obstet Gynecol 2012;24:422–33.
8. Conde-Agudelo A, Papageorghiou AT, Kennedy SH, et al. Novel biomarkers for the prediction of the spontaneous preterm birth phenotype: a systematic review and meta-analysis. BJOG 2011;118:1042–54.
9. Lee L. Likelihood ratios for the prediction of preterm delivery with biomarkers. Acta Obstet Gynecol Scand 2011;90:1189–99.
10. Goldenberg RL, Iams JD, Mercer BM, et al. The Preterm Prediction Study: toward a multiple-marker test for spontaneous preterm birth. Am J Obstet Gynecol 2001;185:643–51.
11. Iams JD, Goldenberg RL, Meis PJ, et al. The length of the cervix and the risk of spontaneous premature delivery. National Institute of Child Health and Human Development Maternal Fetal Medicine Unit Network. N Engl J Med 1996;334:567–72.
12. Smith V, Devane D, Begley CM, et al. A systematic review and quality assessment of systematic reviews of fetal fibronectin and transvaginal length for predicting preterm birth. Eur J Obstet Gynecol Reprod Biol 2007;134:134–42.
13. Abbott DS, Hezelgrave NL, Seed PT, et al. Quantitative fetal fibronectin to predict preterm birth in asymptomatic women at high risk. Obstet Gynecol 2015;125:1168–76.
14. Di Renzo GC. The great obstetrical syndromes. J Matern Fetal Neonatal Med 2009;22:633–5.
15. Brosens I, Pijnenborg R, Vercruysse L, et al. The “Great Obstetrical Syndromes” are associated with disorders of deep placentation. Am J Obstet Gynecol 2011;204:193–201.
16. Committee on Practice Bulletins—Obstetrics, The American College of Obstetricians and Gynecologists. Practice bulletin no. 130: prediction and prevention of preterm birth. Obstet Gynecol 2012;120:964–73.
17. Goldenberg RL, Culhane JF, Iams JD, et al. Epidemiology and causes of preterm birth. Lancet 2008;371:75–84.
18. Honest H, Hyde CJ, Khan KS. Prediction of spontaneous preterm birth. Current Opinion in Obstetrics and Gynecology 2012;24:422–33.
19. Horgan RP, Clancy OH, Myers JE, et al. An overview of proteomic and metabolomic technologies and their application to pregnancy research. BJOG 2009;116:173–61.
20. Wishart DS, Feunang YD, Marcu A, et al. HMDB 4.0: the human metabolome database for 2018. Nucleic Acids Res 2018;46:D608–17.
21. Dettmer K, Hammock BD. Metabolomics—a new exciting field within the "omics" sciences. Environ Health Perspect 2004;112:A396–7.
22. Lucarini F, Morciano L, Rizzo G, et al. Biomarkers for predicting spontaneous preterm birth: an umbrella systematic review. J Matern Fetal Neonatal Med 2018;31:726–34.
23. Romero R, Espinoza J, Gotsch F, et al. The use of high-dimensional biology (genomics, transcriptomics, proteomics, and metabolomics) to understand the preterm parturition syndrome. BJOG 2006;113:118–35.
24. Whiting PF, Rutjes AW, Westwood ME, et al. QUADAS-2: a revised tool for the quality assessment of diagnostic accuracy studies. Ann Intern Med 2011;155:529.
25. Shamseer L, Moher D, Clarke M, et al. Preferred reporting items for systematic review and meta-analysis protocols (PRISMA-P) 2015: elaboration and explanation. BMJ 2015;350:g7647.
26. Zhang A, Sun H, Wang P, et al. Modern analytical techniques in metabolomics analysis. Analyst 2012;137:293–300.
27. McGrath TA, Alaboussi M, Skidmore B, et al. Recommendations for reporting of systematic reviews and meta-analyses of diagnostic test accuracy: a systematic review. Syst Rev 2017;6:194.
28. Kester ADM, Buntinx F. Meta-analysis of ROC Curves. Medical Decision Making 2000;20:430–9.
29. Roselli D. The language of biomedical sciences. Lancet 2016;387:1720–1.