The Zambian Preterm Birth Prevention Study (ZAPPS): Cohort characteristics at enrollment [version 1; referees: 1 approved, 1 approved with reservations]

Marcela C. Castillo¹, Nurain M. Fuseini¹,², Katelyn Rittenhouse¹,², Joan T. Price¹,², Bethany L. Freeman¹, Humphrey Mwape², Jennifer Winston¹, Ntazana Sindano², Courtney Baruch-Gravett³, Benjamin H. Chi¹, Margaret P. Kasaro¹,², James A. Litch³, Jeffrey S.A. Stringer¹,², Bellington Vwalika¹,⁴

¹University of North Carolina at Chapel Hill, Chapel Hill, NC, USA  
²UNC Global Projects Zambia, Lusaka, Zambia  
³Global Alliance for the Prevention of Prematurity and Stillbirth, Seattle, WA, USA  
⁴University of Zambia School of Medicine, Lusaka, Zambia

Abstract

**Background:** Sub-Saharan Africa bears a disproportionate burden of preterm birth and other adverse outcomes. Not only is the background rate of preterm birth higher than in North America and Europe, but many facilities lack essential equipment and personnel resources to care for preterm neonates. A better understanding of the demographic, clinical, and biologic underpinnings of preterm birth is urgently needed to plan interventions and inform new discovery.

**Methods:** The Zambian Preterm Birth Prevention Study (ZAPPS) is a prospective antenatal cohort established at the Women and Newborn Hospital of the University Teaching Hospital (UTH) in Lusaka, Zambia. We recruit pregnant women from the antenatal clinics of district health centers and the UTH for study participation. Women undergo ultrasound examination to determine eligibility by gestational age criteria. Enrolled participants receive routine antenatal and postnatal care, lab testing, midtrimester cervical length measurement, serial fetal growth monitoring and careful assessment of birth outcomes.

**Results:** Between August 2015 and September 2017, we screened 1784 women, of whom 1450 (81.2%) met inclusion criteria and were enrolled. The median age at enrollment of study participants is 27 years (IQR 23–32). Participants are enrolled at a median gestational age of 16 weeks (IQR 13–18). Among all parous participants (N=866; 64%), 21% (N=182) reported a prior miscarriage, 49% (N=424) reported a prior preterm birth, and 13% (N=116) reported a prior stillbirth. The HIV seroprevalence in our cohort is 24%.

**Discussion:** We have established a large antenatal cohort to characterize the
epidemiological and biological determinants of adverse birth outcomes in Lusaka, Zambia. Findings from this cohort will help guide future studies, clinical care, and policy in the prevention and treatment of adverse birth outcomes.

Keywords
Preterm Birth, Africa, Cohort
Introduction
Preterm birth is a global challenge impacting both developed and developing countries\(^1\). It contributes to approximately 35% of neonatal and 75% of perinatal mortality each year\(^4\). Further, preterm infants who survive are at an elevated risk of long-term respiratory, cardiovascular, gastrointestinal, and neurodevelopmental morbidities. These complications may affect subsequent health, growth, psychosocial functioning, and even economic capacity of these individuals\(^5\).

The greatest burden of mortality and morbidity from preterm births occur in low- and middle-income countries (LMICs)\(^6\). Of an estimated 14.9 million preterm births globally each year, 13.6 million (91%) occur in LMICs\(^7\). Preterm birth rates are as low as 5% in some European countries and as high as 18% in some African countries\(^8\), precisely where the resources to prevent preterm birth and manage preterm infants are least developed. In Zambia, for example, the preterm birth rate is estimated to be 13%\(^9\). Each year there are 77,600 preterm births and 6,800 infant deaths due to preterm birth complications\(^9\).

The burden of maternal HIV infection is also high in many LMIC settings, where it has been associated with a 50% increased risk of preterm birth\(^10\). Although the increasing availability of maternal antiretroviral therapy has led to dramatic reductions in pediatric HIV incidence\(^11\), it does not seem to reduce HIV-attributable preterm birth in this population. In fact, antiretroviral drug exposure may in fact increase the risk of preterm birth among some HIV-infected gravidas\(^12\)-\(^14\).

Preterm birth can result from many different etiological entities. Approximately one-third of preterm deliveries are indicated because of pre-eclampsia, hemorrhage, abnormal placentation, intra-uterine growth restriction, oligohydramnios, or multi-fetal gestation. Spontaneous preterm labor is implicated in about 40% of preterm births; another 25% are related to preterm premature rupture of the membranes\(^15\)-\(^17\). The underlying causes of spontaneous preterm birth in HIV-infected and HIV-uninfected populations are not well understood. Although several maternal and newborn interventions (e.g. antenatal corticosteroids, neonatal resuscitation, and kangaroo mother care) can reduce the complications of preterm birth, prevention is key. Much remains to be discovered about the risk factors, causes, and pathophysiology of preterm delivery, and how to prevent its occurrence.

Methods
Study design
The Zambian Preterm Birth Prevention Study (ZAPPS) aims to establish a well-characterized pregnancy cohort to better understand the risk factors associated with preterm birth and other adverse birth outcomes in a LMIC setting. The cohort was established to be a local resource to the University of Zambia School of Medicine and to contribute to general scientific knowledge around the biology of pregnancy and parturition.

ZAPPS enrolls pregnant women at the Women and Newborn Hospital of the University Teaching Hospital (UTH) in Lusaka, Zambia into a prospective antenatal cohort. The UTH is the province’s only tertiary referral center, serving a primary catchment population of approximately 2 million people. The Women and Newborn Hospital receives referrals for high-risk pregnancies, including those with complex medical histories, history of prior preterm birth, stillbirth, or pregnancy loss, and has a very busy labor ward with approximately 18,000 deliveries per year\(^18\). Study participants are recruited from the UTH and five nearby high-volume Lusaka district health clinics. We established this cohort with the aim to better characterize demographic determinants, biomedical causes, and underlying pathophysiologic mechanisms associated with adverse birth outcomes in Lusaka, Zambia. Through the Global Alliance to Prevent Prematurity and Stillbirth, we collaborate with a consortium of international scientists, many of whom are also working in LMIC countries, in order to advance our understanding of the causes of preterm delivery. This study was designed in accordance with the Strengthening The Reporting of Observational Studies in Epidemiology (STROBE) guidelines\(^19\).

Study participants
Pregnant women who meet the following criteria are eligible for enrollment in ZAPPS: (1) 18 years of age or older; (2) viable intrauterine singleton or twin pregnancy; (3) presentation to antenatal care prior to 20 weeks’ gestation if HIV-uninfected or 24 weeks’ if HIV-infected; (4) residing within Lusaka with no plans to relocate during the study follow-up period; (4) willing to provide written, informed consent; (5) willing to allow participation of their infant(s) in the study; (6) willing to be followed up at home for birth outcomes if necessary. Initially, all pregnant women with a gestational age \(\geq 20\) weeks by a standard algorithm\(^20\) were excluded; however, this criterion was extended to \(\geq 24\) weeks for HIV-infected women following a protocol amendment in July 2016 to align this group with related, ongoing clinical trials at this site (NCT03297216, NCT02970552).

Primary objective
Our primary objective in this study is to establish a well-characterized cohort of pregnant women and their infants – and an accompanying specimen biorepository – with follow-up through delivery and the postpartum period.

Study procedures
Potential participants are identified at an early antenatal visit. Community educators approach participants who may be preliminarily eligible by gestational age criteria based on reported last menstrual period and fundal height. Potentially eligible women are escorted to the UTH study clinic. After an information session, a sonographer performs an ultrasound examination to determine fetal viability and gestational age estimation either by crown rump length (if \(<14\) weeks)\(^19\) or fetal biometry (if \(\geq 14\) weeks) measurements\(^22\). Women deemed eligible for study participation and choosing to participate are administered an informed consent in the language of their
choice: English, Nyanja, or Bemba. While screening and enrollment procedures may occur on the same day, women could return on a subsequent day for enrollment to allow time to consider the risks and benefits of study participation and to discuss the study with their family.

Clinical care and follow-up
Study participants receive routine antenatal care at the ZAPPS study clinic at the UTH, with visits scheduled at enrollment, 24 weeks, 32 weeks, and 36 weeks of gestation, according to standard of care in Zambia. Additionally, women are asked to return to the clinic for a postpartum visit, typically 6 weeks after delivery (Table 1).

After enrollment, participants return for cervical length evaluation between 20 and 24 weeks’ gestation. Women with a short cervix by transvaginal ultrasound, defined as <2.5 cm², attend additional visits scheduled at 28 and 32 weeks’ gestation for repeat cervical length ultrasounds and are referred to a study physician at the UTH for further counseling and follow-up. All participants undergo an additional fetal biometry ultrasound, performed at 32 weeks’ gestation. Each study sonographer is trained using curricula adapted from the INTERGROWTH-21,24 and Cervical Length Education and Review (CLEaR) program for cervical length measurements. All biometry parameters are measured twice and then averaged. Cervical length – measured three times over a period of 3 to 5 minutes – is first measured by transabdominal ultrasound; those whose transabdominal cervical length is <3.5cm or not measurable then undergo transvaginal measurement25,26.

At each antenatal care visit, study nurses perform a vital sign assessment and a physical exam. We use point-of-care tests for HIV, anemia, malaria, syphilis, and urinary tract infection, and provide tetanus toxoid injection(s), iron, folate, malaria intermittent preventive treatment, and de-worming treatment in accordance with local standards of care. Participants with pre-existing or new HIV diagnoses are counseled and referred to appropriate antiretroviral therapy and prevention of mother-to-child transmission services. At the postpartum visit, study nurses assess maternal and infant interval complications, perform maternal and infant physical exams, assess infant feeding and general well-being, and provide health education counselling.

Throughout the study, study nurses assess participants’ past medical and obstetrical history and current pregnancy signs and symptoms. Study staff carefully screen participants at each study visit for the presence of adverse or serious adverse events. Participants are referred to the appropriate higher level care provider at the UTH for any adverse events identified that require medical care beyond the scope of the study nurses’ practice.

To maximize retention, locator information on all participants is collected at screening and reviewed at each subsequent encounter. All participants are informed during the consent process that their locator sources will be used to contact them if they do not attend their scheduled study visits. Missed visits are identified by an electronic database that tracks expected and actual visits. If a participant misses a scheduled visit, study staff follow standardized procedures to attempt to contact the participant through the following mechanisms: (1) phone contact with the participant directly, (2) phone contact with other contacts provided on the participant’s locator information, and (3) home visits.

Data collection and management
Clinical data: After enrollment, study staff collect medical, antenatal, and HIV history data (as applicable) through interviewer-administered questionnaires and review of participants’ medical records. At the time of delivery, or at first contact postpartum, detailed information is collected about the clinical course of the participant’s delivery and delivery outcomes for both the mother and infant(s). This allows clinical phenotyping of all adverse birth outcomes. Shortly after delivery and prior to hospital discharge, the study team documents assessment of size, growth, and physical and neuromuscular maturity. If the infant requires admission to the neonatal intensive care unit, the newborn assessment is done after the infant is deemed stable by the pediatrician.

Biological specimen collection: Trained study staff collect maternal specimens at enrollment, 24-week, and 32-week visits, as well as at delivery (Table 1) following approved standard operating procedures. All specimens are stored and transferred in temperature-controlled containers to the on-site lab by clinic staff within two hours of collection. Study lab staff process all specimens according to assay manufacturers’ instructions, analyzing some specimens immediately per standard antenatal care guidelines and storing others for later study-related analyses. HIV-1 plasma viral loads are performed for participants identified as HIV-infected at enrollment. Lab staff follow strict quality protocols for maternal blood processing to produce aliquots of whole blood, serum, plasma, anduffy coat for storage in barcoded cryotubes. Trained study nurses collect maternal specimens pre-delivery as well as placenta and cord blood specimens immediately following delivery. The UTH serves as the primary biorepository for stored specimens, with redundancy at a central project repository in Seattle, Washington and at the University of North Carolina at Chapel Hill in Chapel Hill, NC.

Ethical considerations
The ZAPPS protocol was developed in consultation with a local community advisory board to ensure study procedures are acceptable in the communities from which participants would be recruited. The study and its protocol revisions undergo continuing ethical review by the relevant research ethics authorities at the University of Zambia School of Medicine (Reference number: 016-04-14) and the University of North Carolina School of Medicine (Study number: 14-2113). Participation in all study
<table>
<thead>
<tr>
<th>Table 1. Schedule of events, demographic and clinical data collected among all participants in ZAPPS cohort.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gestational age (weeks)</strong></td>
</tr>
<tr>
<td><strong>ADMINISTRATIVE/REGULATORY PROCEDURES</strong></td>
</tr>
<tr>
<td>Informed consent</td>
</tr>
<tr>
<td>Collection/review of locator info</td>
</tr>
<tr>
<td><strong>COLLECTION OF DEMOGRAPHIC VARIABLES</strong></td>
</tr>
<tr>
<td>Age, education, socioeconomic status</td>
</tr>
<tr>
<td>Substance use</td>
</tr>
<tr>
<td>Marital status</td>
</tr>
<tr>
<td>Pregnancy intention</td>
</tr>
<tr>
<td>Sexual health</td>
</tr>
<tr>
<td>Vaginal practices</td>
</tr>
<tr>
<td>Intimate partner violence screening</td>
</tr>
<tr>
<td>Nutritional assessment</td>
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<tr>
<td>Maternal depression screen</td>
</tr>
<tr>
<td><strong>OBSTETRICAL ULTRASOUND PROCEDURES</strong></td>
</tr>
<tr>
<td>Dating biometry ultrasound</td>
</tr>
<tr>
<td>Fetal biometry ultrasound</td>
</tr>
<tr>
<td>Cervical length ultrasound</td>
</tr>
<tr>
<td><strong>MATERNAL CLINICAL HISTORY AND PHYSICAL EXAM</strong></td>
</tr>
<tr>
<td>Obstetrical history</td>
</tr>
<tr>
<td>Medical history</td>
</tr>
<tr>
<td>Maternal height &amp; weight</td>
</tr>
<tr>
<td>Maternal mid-upper arm circumference</td>
</tr>
<tr>
<td>Maternal vital signs</td>
</tr>
<tr>
<td>Maternal physical exam</td>
</tr>
<tr>
<td>Fetal heart rate</td>
</tr>
<tr>
<td>Fundal height</td>
</tr>
<tr>
<td>Fetal lie</td>
</tr>
<tr>
<td><strong>INFANT CLINICAL HISTORY AND PHYSICAL EXAM</strong></td>
</tr>
<tr>
<td>Neonatal physical exam</td>
</tr>
<tr>
<td>Neonatal vital status / APGAR</td>
</tr>
<tr>
<td>Newborn assessment</td>
</tr>
<tr>
<td>Infant physical exam</td>
</tr>
<tr>
<td>Infant feeding status assessment</td>
</tr>
<tr>
<td>Infant HIV diagnostic assessment (if exposed)</td>
</tr>
<tr>
<td><strong>LABORATORY PROCEDURES</strong></td>
</tr>
<tr>
<td>Maternal HIV (rapid EIA)</td>
</tr>
<tr>
<td>Maternal syphilis (RPR)</td>
</tr>
<tr>
<td>Maternal malaria</td>
</tr>
<tr>
<td>Maternal hemoglobin (hemocue)</td>
</tr>
<tr>
<td>Maternal urinalysis (&amp; culture if +)</td>
</tr>
<tr>
<td><strong>SPECIMEN COLLECTION FOR STORAGE</strong></td>
</tr>
<tr>
<td>Vaginal ± rectal swab storage</td>
</tr>
<tr>
<td>Blood storage</td>
</tr>
<tr>
<td>Urine storage</td>
</tr>
<tr>
<td>Placenta, membranes, cord histopathology, and storage</td>
</tr>
<tr>
<td>Infant blood sample via heel prick</td>
</tr>
</tbody>
</table>

<sup>†</sup> Extended to <24 weeks for HIV-infected in July 2016

<sup>‡</sup> Additional visits for participants with short cervix

Birth weight, birth length, head circumference, foot length, physical and neuromuscular maturity
activities is voluntary, and each participant provides written, informed consent prior to enrollment.

To address the minimal risks associated with participation in this non-interventional study, all study personnel have been trained on standard operating procedures to protect participant privacy and confidentiality. Staff receive protection of human research participants training prior to conducting any study activities and every two years thereafter. Key research staff members complete Good Clinical Practice or Good Clinical Laboratory Practice training, as applicable. All study-related and unrelated adverse events and social harms are graded using the National Institute of Health’s Division of AIDS Table for Grading the Severity of Adult and Pediatric Adverse Events. All adverse events are reported to regulatory authorities, according to their individual guidelines.

All participants in this study may benefit from enhanced health education and close clinical monitoring. The knowledge generated from this observational study regarding maternal risk factors and neonatal outcomes of preterm birth are expected to outweigh the risks of participation. Conclusions drawn from this knowledge may inform future clinical trials on the prevention of adverse birth outcomes in low-resource settings, which may in turn enable policymakers worldwide to make informed decisions regarding effective interventions for the improvement of maternal-neonatal health.

**Results**

**Characteristics of cohort at enrollment**

Between August 2015 and September 2017, 1784 women were recruited and screened from local antenatal clinics by ZAPPS study staff. Among them, 1450 (81.3%) met inclusion criteria and were enrolled. Of the 334 not enrolled, 274 (82%) were at advanced gestational age on ultrasound, 7 (2%) were less than 18 years old, 12 (4%) were unwilling to provide informed consent, 5 (1%) were unwilling to remain in the study area, and 36 (11%) were not enrolled for other reasons.

The median age of participants in the cohort is 27 years (IQR 23–32) (Table 2). Most women (n=1201 of 1435; 84%) are married or cohabiting with their partner, and have been pregnant at least once in the past (n=866 of 1352, 64%). Among parous participants, 21% (n=182) reported a prior miscarriage, 49% (n=424) reported a prior preterm birth, and 13% (N=116) reported a prior stillbirth. Participants are enrolled at a median gestational age of 16 weeks (IQR 13–18); 421 of 1428 (29%) enrolled prior to 14 weeks' gestational age.

### Table 2. Baseline characteristics of ZAPPS cohort, N=1450.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>N</th>
<th>Value* % or Median (IQR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>1406</td>
<td>27 (23;32)</td>
</tr>
<tr>
<td>&lt;20</td>
<td>111</td>
<td>7.9</td>
</tr>
<tr>
<td>20–34</td>
<td>1113</td>
<td>79.2</td>
</tr>
<tr>
<td>≥35</td>
<td>182</td>
<td>12.9</td>
</tr>
<tr>
<td>Missing</td>
<td>44</td>
<td>-</td>
</tr>
<tr>
<td>Marital status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not married and not cohabiting</td>
<td>234</td>
<td>16.3</td>
</tr>
<tr>
<td>Married or cohabiting</td>
<td>1201</td>
<td>83.7</td>
</tr>
<tr>
<td>Missing</td>
<td>15</td>
<td>-</td>
</tr>
<tr>
<td>Education</td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>26</td>
<td>1.8</td>
</tr>
<tr>
<td>&lt;8 years</td>
<td>243</td>
<td>17.0</td>
</tr>
<tr>
<td>8-9 years</td>
<td>328</td>
<td>22.9</td>
</tr>
<tr>
<td>10-12 years</td>
<td>653</td>
<td>45.5</td>
</tr>
<tr>
<td>≥12 years</td>
<td>184</td>
<td>12.8</td>
</tr>
<tr>
<td>Missing</td>
<td>16</td>
<td>-</td>
</tr>
<tr>
<td>Source of drinking water</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Piped</td>
<td>1339</td>
<td>93.3</td>
</tr>
<tr>
<td>Other</td>
<td>96</td>
<td>6.7</td>
</tr>
<tr>
<td>Missing</td>
<td>15</td>
<td>-</td>
</tr>
<tr>
<td>Toilet facilities in household</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flush or Pour</td>
<td>762</td>
<td>53.1</td>
</tr>
<tr>
<td>Pit or Latrine</td>
<td>672</td>
<td>46.8</td>
</tr>
<tr>
<td>Other</td>
<td>2</td>
<td>0.1</td>
</tr>
<tr>
<td>Missing</td>
<td>14</td>
<td>-</td>
</tr>
<tr>
<td>Floor material in home</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Natural/rudimentary</td>
<td>138</td>
<td>9.6</td>
</tr>
<tr>
<td>Finished</td>
<td>1298</td>
<td>90.4</td>
</tr>
<tr>
<td>Missing</td>
<td>14</td>
<td>-</td>
</tr>
<tr>
<td>BMI, kg/m2</td>
<td>1371</td>
<td>23.6 (21.2;27.2)</td>
</tr>
<tr>
<td>&lt;18.5</td>
<td>72</td>
<td>5.3</td>
</tr>
<tr>
<td>18.5–30.0</td>
<td>1104</td>
<td>80.5</td>
</tr>
<tr>
<td>&gt;30.0</td>
<td>195</td>
<td>14.2</td>
</tr>
<tr>
<td>Missing</td>
<td>79</td>
<td>-</td>
</tr>
<tr>
<td>GA at enrollment, weeks</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;14</td>
<td>421</td>
<td>29</td>
</tr>
<tr>
<td>≥14</td>
<td>1007</td>
<td>69.5</td>
</tr>
<tr>
<td>Missing</td>
<td>22</td>
<td>-</td>
</tr>
</tbody>
</table>
The baseline HIV seroprevalence in our cohort is 24% (n=350 of 1447), of whom 60% (n=205 of 340) had undetectable viral load. Nearly 4% (n=52 of 1424) of participants had elevated blood pressure (≥140/90) at enrollment; 5% (n=69 of 1372) had a urinalysis consistent with bacteriuria or urinary tract infection (1+ leukocyte esterase and/or nitrates). Syphilis was prevalent in 5% (n=70 of 1343) of our cohort at baseline. Malaria is uncommon in our cohort: 5 of 1148 participants (0.4%) tested positive for malaria by rapid test. 14% (n=141 of 1026) were anemic (Hgb <10.5mg/dL) at baseline.

**Discussion**

The underlying pathological processes responsible for activation of the common parturition pathway in preterm labor, preterm prelabor rupture of membranes, and preterm delivery are incompletely understood. A comprehensive investigation of these processes could help define clinical signs and biological markers of high-risk pregnancies and allow the development and application of early preventive interventions targeted to those women at highest risk. Our cohort study will describe incidence, predictors, and potential confounders to the parturition pathway in HIV-infected and HIV-uninfected term and preterm deliveries, in accordance with the International Preterm Birth Collaborative Pathways and Systems Biology Working Group.

Analyses of the ZAPPS cohort data will investigate determinants of the following adverse birth outcomes, both individually and in composite: preterm birth (delivery <37 weeks), very preterm birth (delivery <34 weeks), stillbirth, low birthweight (<2500g), very low birthweight (<1500g), small for gestational age (<10%ile), and very small for gestational age (<3%ile). We will define birthweight-for-age according to INTERGROWTH-21 standards. We will distinguish spontaneous preterm deliveries from provider-indicated preterm deliveries and investigate the prevalence and distribution of preterm phenotypes, both within our cohort and relative to other studies. Specimens stored at our UTH laboratory will be used for study-related analyses to identify inflammatory markers (e.g., chemokines, cytokines), microbiome community states, metabolic analytes, proteins, hormones, transcripts, and various infectious factors that may be predict adverse birth outcomes. As nearly one-fourth of our cohort was HIV-infected at enrollment, we will conduct specific analyses within this population to better understand the effect of both HIV and antiretroviral therapy on adverse birth outcomes.

Gestational age can be estimated by patient report of the last normal menstrual period (LMP), ultrasound biometry, or a combination of the two. Our experience in Zambia is that the LMP is an imprecise measure that artificially inflates the actual rate of preterm birth in the population, and that it may in fact introduce bias. We chose to establish gestational age by ultrasound alone in this cohort.

The strength of ZAPPS is found in its size (nearly 1500 participants enrolled to date) and its design (a prospective antenatal cohort enrolling in early pregnancy). We note that our cohort is at risk of attrition, a well-known challenge of antenatal cohorts, as well as biases of selective participation contributing to a cohort not fully representative of the general population. We enroll pregnant women in the nation’s capital of Lusaka, and we have specifically prioritized the enrollment of HIV-infected women, relaxing the eligibility criteria for gestational age at enrollment for this population. This over-representation of...
HIV-infected participants in our cohort will enhance our ability to investigate epidemiologic and mechanistic associations of HIV and preterm birth.

In summary, we have established a well-characterized antenatal cohort in Lusaka, Zambia that benefits from ultrasound gestational age dating, longitudinal clinical assessments, biological specimen collection and storage, and careful classification of birth and neonatal outcomes (including phenotyping of all preterm births and stillbirths). The knowledge gained from this study has the potential to drive future research in preterm birth and other adverse birth outcomes, to inform the development of novel preventive therapies and treatments, and to influence clinical care and health policy worldwide.

Collaboration
The ZAPPS study is part of the Preventing Preterm Birth Initiative of the Global Alliance to Prevent Prematurity and Stillbirth (GAPPS). The Zambia cohort is co-led by the University of Zambia and the University of North Carolina at Chapel Hill. Study findings will be made available through appropriate local channels, including academic and public health research symposia. Our primary purpose is as a shared resource and we invite collaborators with high-impact ideas to apply for access to data and stored specimens from the ZAPPS study as well as other sites in the GAPPS biorepository network. Potential collaborators are invited to contact GAPPS directly (info@gapps.org) or the ZAPPS principal investigators: Jeffrey Stringer (jeffrey_stringer@med.unc.edu) and Bellington Vwalika (bvwalika@unza.zm).

Data availability
De-identified individual patient data underlying Table 2 are available on Open Science Framework: http://doi.org/10.17605/OSF.IO/UNE9Y

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

Grant information
Bill and Melinda Gates Foundation grant to the Global Alliance to Prevent Prematurity and Stillbirth (Seattle Children’s Hospital/GAPPS 13008/OPP1033514). Additional support was provided by the US National Institutes of Health through the UNC Center for AIDS Research (P30 AI50410) and trainee/mentor support: T32 HD075731 (MCC, NMF, JTP), K01 TW010857 (JTP), D43 TW009340 (KR), and K24 AI120796 (BHC).

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Acknowledgments
The study protocol is registered at ClinicalTrials.gov, identifier: NCT02738892.

References


Data Source
Open Peer Review

Current Referee Status: ✅ ?

Version 1

Referee Report 24 May 2018

https://doi.org/10.21956/gatesopenres.13888.r26413

Fyezah Jehan  
Department of Pediatrics and Child Health, Aga Khan University, Karachi, Pakistan

Overall an important report of a biorepository cohort in LMIC, in a high prematurity setting defining baseline characteristics of the cohort at enrollment. Some recommendations for improvement:

Abstract: Needs to be realigned with the article text. Methods incomplete, all salient points need to be mentioned eg. GA at eligibility, samples being stored in biorepository. Similarly, what outcome information is being collected. Conclusion mentions adverse birth outcomes although focus in the introduction has been on preterm.

Article:

Introduction: Focus of discussion has been on prematurity its risk factors and causes, while the title, methods, objectives are related to establishment of a bio repository. It isn't entirely clear if this is a description of the biorepository or a description of cohort characteristics of the women in the biorepository or both. A clearer enunciation of the objective needs to be made. Adverse outcomes to be studied are detailed in the discussion session so I would suggest to either bring that detail into the methods, in order to describe the biorepository completely.

Methods:
Clinical care and followup/ Data collection and management: Overall lacks specificity in discussion for e.g. what particular medical history and exam is done? How is size, neuromuscular maturity assessed? Is there a window of collection? Who does it? What's the training? What is the quality control? How are the specimens stored? What particular tests are anticipated? What is meant by temperature-controlled? 'analyzing some specimens immediately per standard antenatal care guidelines and storing others for later study-related analyses"can be unpacked.

Results:
There is some redundancy in the results text when looking at the table and vice versa. Not sure how the authors can handle this. In conformity with data and specimen collection as described in methods, advised to also detail the specimens collected so far?

Table 2:
Education- too many categories. Can collapse.
GA missing on 22 women- how were they included?
BMI-Typo, should be 13.5 not 13-5

Discussion: Again can be realigned with the results presentation.
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?
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:** I am the PI of the AMANHI-ACT Cohort and Biorepository in Pakistan

**Referee Expertise:** Maternal newborn health, community, biorepository, cohort studies, clinical trials, infectious diseases

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Author Response 12 Nov 2018**

**Jeffrey S. A. Stringer,** University of North Carolina, USA

**Fyezah Jehan,** Department of Pediatrics and Child Health, Aga Khan University, Karachi, Pakistan

**Approved with Reservations**

Overall an important report of a biorepository cohort in LMIC, in a high prematurity setting defining baseline characteristics of the cohort at enrollment. Some recommendations for improvement:

**Abstract:** Needs to be realigned with the article text. Methods incomplete, all salient points need to be mentioned eg. GA at eligibility, samples being stored in biorepository. Similarly, what outcome information is being collected. Conclusion mentions adverse birth outcomes although focus in the introduction has been on preterm.

Thank you for noting these inconsistencies. We have expanded our description of assessment of birth outcomes to include not only gestational age at delivery, but also “neonatal birthweight, vital status, and sex” (Abstract, lines 17-18). We have added the gestational age criteria for enrollment (lines 14-15), as well as a description of sample collection to the methods section of the abstract (lines 18-20). Finally, we now refer to “other adverse outcomes” to the final sentence of the
background of the abstract (line 7).

**Article:**

**Introduction:** Focus of discussion has been on prematurity its risk factors and causes, while the title, methods, objectives are related to establishment of a bio repository. It isn't entirely clear if this is a description of the bio repository or a description of cohort characteristics of the women in the biorepository or both. A clearer enunciation of the objective needs to be made. Adverse outcomes to be studied are detailed in the discussion session so I would suggest to either bring that detail into the methods, in order to describe the biorepository completely.

The purpose of this cohort description is to outline baseline characteristics of the ZAPPs Phase 1 cohort. We mention the biorepository, but that is not the primary aim of this paper. We have clarified the role of the biorepository throughout the abstract and manuscript, but note that the purpose of this paper is primarily to describe the clinical characteristics of our population.

The second paragraph in the discussion section (lines 268-280) details the outcomes to be studied in future analyses. We think it would be confusing to bring that detail into the methods of this paper as these analyses have not yet been conducted and we have not included any outcomes in this baseline cohort description. We have attempted to clarify this in said paragraph.

**Methods:**

Clinical care and followup/ Data collection and management: Overall lacks specificity in discussion for e.g. what particular medical history and exam is done? How is size, neuromuscular maturity assessed? Is there a window of collection? Who does it? What's the training? What is the quality control? How are the specimens stored? What particular tests are anticipated? What is meant by temperature-controlled? 'analyzing some specimens immediately per standard antenatal care guidelines and storing others for later study-related analyses"can be unpacked.

We have added / clarified the following:

1. Lines 146-149: “At each antenatal care visit, study nurses perform a vital sign assessment and a physical exam, which includes maternal height and weight; mid-upper arm circumference measurement; fundal height measurement; assessment for pallor, edema, and abdominal tenderness; fetal heart rate assessment; and cervical exam as clinically indicated.”
2. Lines 161-163: “Participants are asked specifically if they have ever been diagnosed with high blood pressure, heart disease, diabetes, HIV/AIDS, tuberculosis, or any other chronic illness.”
3. Lines 180-183: “Shortly after delivery and prior to hospital discharge, the study team documents assessment of infant vital signs, weight, length, head circumference, complete physical exam, and of neuromuscular maturity via the New Ballard Score.”
4. Lines 187-189: “While the 24- and 32-week study visits are scheduled according to gestational age, in the event that a participant misses her appointment, specimens may be collected as soon as possible once she returns to clinic.”
5. Lines 185-187: “Trained study nurses collect maternal specimens...”
6. Lines 186-187: “following approved standard operating procedures to ensure quality and uniformity.”
7. Lines 275-280: Possible protocol-related tests anticipated are described in the discussion section: “Specimens stored at our UTH laboratory will be used for study-related analyses to
identify inflammatory markers (e.g., chemokines, cytokines), microbiome community states, metabolic analytes, proteins, hormones, transcripts, and various infectious factors that may be predict adverse birth outcomes.”

8. Lines 191-204: “All specimens are transferred in insulated containers with continuous temperature monitoring to the on-site lab” … “All specimens are stored at -80 C in temperature-controlled freezer systems equipped with continuous temperature monitoring and text message notification of temperature deviations.”

9. Line 199: We have added a reference to Table 1, which delineates exactly which tests are performed immediately and we discuss which types of study-related testing may be performed in the future in the Discussion as outlined above.

Results:
There is some redundancy in the results text when looking at the table and vice versa. Not sure how the authors can handle this.

We prefer to retain some redundancy between Table 1 and the Results section. Many journals require this (e.g., JAMA). The description of key baseline characteristics of our cohort in the Results section is not meant to be exhaustive, but rather to highlight key data that appear in Table 1.

In conformity with data and specimen collection as described in methods, advised to also detail the specimens collected so far?

While specimens have been collected, given this is a baseline paper of cohort characteristics at enrollment, we have not presented data on the total numbers of specimens that have been collected to date. This will be addressed in a future publication.

Table 2:
Education- too many categories. Can collapse.
We have collapsed this to: “None”, “0-12 years”, and “≥12 years”

GA missing on 22 women- how were they included?
Thank you for noting this. After additional data cleaning since our first submission of this manuscript, we have now been able to assign entry gestational age for each cohort participant.

BMI-Typo, should be 13.5 not 13-5
We have corrected this to “18.5” from “18-5”

Discussion: Again can be realigned with the results presentation.
Thank you. We feel that as a result of these suggested revisions, the discussion and results section are now more closely aligned.

Competing Interests: No competing interests were disclosed.
Thank you for giving me the opportunity to review the research article entitled “The Zambian Preterm Birth Prevention Study (ZAPPS: Cohort characteristics at enrollment)”

Overall the study is important to understand the dynamics of preterm birth in a sub-saharan developing country, with intent to plan interventions.

I have the following comments: –

Abstract:
Methods – should read as ‘the Zambian Preterm Birth Prevention study (ZAPPS) is an observational study, of a prospective antenatal cohort ………’
Results- this cohort is 1450 participants and is not clear is this the entire cohort or a subset as the protocol mentions the enrollment to be estimated at 4000, so is recruitment still ongoing? This needs to be clarified.
Discussion – also add the primary objective of the ZAPPS study, as stated below

Page 3
Primary objective – should also include the primary objective of the study as stated in the protocol
Primary outcome measure – Rate of preterm births per 100 person years, incidence rate of preterm birth (delivery <37 weeks’ gestation)

Page 4 Clinical care and follow up
It is stated that after enrollment women return for cervical length evaluation between 20 and 24 weeks and those with a short cervix have additional visits at 28 and 32- in table 1 the ultrasound is at 34 and not 32 weeks. It is important to mention if this is standard of care for all patients at this site or only for those with previous preterm birth. Page 6 mentions that this is a non-interventional study.
No mention is made of the maternal depression screen at 24 and 42 weeks.

Page 4 Biological specimen collection
Study nurses collected maternal specimens at enrollment –and these were point of care tests for HIV, anemia, syphilis and UTI.
The results section has a reasonably large number of missing information on some of the parameters.
Page 5 table 1
What were the vaginal and rectal swab tests for?

Page 6
Results section
Table 2- the missing numbers of participates with data on Syphilis is 107, Haemoglobin at enrollment 424 and malaria is 302, this needs to be explained by the authors. Patients with malaria, low haemoglobin and syphilis are at risk for preterm birth, so missing data will influence results for the ZAPPS study.
This should be discussed by the authors as a study limitation if these figures are correct.

This comment does not influence this paper as it for those recruited up to Sept 2017.

Of note is that the protocol for ZAPPS has been amended in Feb 2018 to exclude women with HIV. So eventually will there be separate analysis for the women who are included in this manuscript vs those recruited later?

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?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Not applicable

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.

Referee Expertise: Obstetrician/ gynaecologist with interest in Maternal fetal medicine

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

Author Response 12 Nov 2018

Jeffrey S. A. Stringer, University of North Carolina, USA

Zahida Qureshi, Department of Obstetrics and Gynaecology, School of Medicine, University of Nairobi, Nairobi, Kenya

Approved

Thank you for giving me the opportunity to review the research article entitled “The Zambian Preterm Birth Prevention Study (ZAPPS: Cohort characteristics at enrollment)”

Overall the study is important to understand the dynamics of preterm birth in a sub-saharan developing country, with intent to plan interventions.

I have the following comments: -

Abstract:
Methods – should read as ‘the Zambian Preterm Birth Prevention study (ZAPPS) is an observational study, of a prospective antenatal cohort ……….’
This suggested edit has been made (line 10).

Results-this cohort is 1450 participants and is not clear is this the entire cohort or a subset as the protocol mentions the enrollment to be estimated at 4000, so is recruitment still ongoing? This needs to be clarified.
Recruitment is indeed ongoing, but under a slightly revised protocol. This is a description of the cohort established under the original protocol only. The original estimated enrollment of 4000 has been amended to 2000 on the ClinicalTrials.gov website.
Discussion—also add the primary objective of the ZAPPS study, as stated below
Our primary objective in the protocol reads as follows: “to establish a cohort of 2,000 pregnant women and their infants, following them through delivery and up to 42 days postpartum.” We have revised the language in the Discussion of the abstract to reflect that original wording (lines 31-32).

Page 3
Primary objective—should also include the primary objective of the study as stated in the protocol
Our primary objective in the protocol reads as follows: “to establish a cohort of 2,000 pregnant women and their infants, following them through delivery and up to 42 days postpartum. We will recruit and enroll 2,000 women and their newborn infants in Zambia.” We have revised the language in our “Primary objective” section to reflect this wording (lines 106-110).

Primary outcome measure—Rate of preterm births per 100 person years, incidence rate of preterm birth (delivery <37 weeks' gestation)
Thank you for noting this inconsistency. The primary outcome as stated in the most recent protocol is as follows:
“The primary outcome is preterm birth, defined as birth prior to 37 weeks completed gestation. The secondary outcomes are (1) delivery prior to 34 weeks completed gestation and (2) low birth weight, defined as less than 2500 grams.”
We have added an “Outcomes” section to the Methods of this paper to address this omission (lines 111-118).

Page 4 Clinical care and follow up
It is stated that after enrollment women return for cervical length evaluation between 20 and 24 weeks and those with a short cervix have additional visits at 28 and 32 weeks. It is important to mention if this is standard of care for all patients at this site or only for those with previous preterm birth. Page 6 mentions that this is a non-interventional study.
We have revised the text to the following: “After enrollment, all participants return for universal cervical length evaluation between 20 and 24 weeks’ gestation. Those with a short cervix by transvaginal ultrasound, defined as <2.5 cm, then attend additional visits scheduled at 28 and 32 weeks’ gestation for repeat cervical length ultrasounds and are referred to a study physician at the UTH for further counseling and follow-up” (lines 135-139). The additional visits at 28 and 34 weeks’ for cervical length ultrasound are only for those women with short cervix. This is explained also in the footnote to Table 1. Biometry ultrasound at 32 weeks is performed for all participants.

No mention is made of the maternal depression screen at 24 and 42 weeks.
Thank you for pointing this out. The following sentence has been added to the section on Clinical care & follow-up:
“The Edinburgh Postnatal Depression Screen is self-administered by participants at 24 weeks and again at the postpartum visit. Participants who screen positive are referred to outpatient psychiatric care” (lines 157-159). In addition, “Maternal depression screen” appears in Table 1 at 24 weeks and 42 days postpartum.

Page 4 Biological specimen collection
Study nurses collected maternal specimens at enrollment—and these were point of care tests for HIV, anemia, syphilis and UTI.
The results section has a reasonably large number of missing information on some of the parameters.

Since our study is integrated with routine care at the clinical sites, for several cases—particularly in the early conduct of the study—we did not repeat routine tests previously performed at government clinics and failed to document these results on our study forms. This is a limitation of the study that we have described in our Discussion (lines 295-299). We also note that it has been rectified in Phase 2 of the cohort (line 299).

**Page 5 table 1**

What were the vaginal and rectal swab tests for?

Vaginal and rectal swabs are collected for specimen storage for future protocol-related testing. No real-time tests are currently being performed on these swabs. We have clarified this in Table 1.

**Page 6**

**Results section**

Table 2- the missing numbers of participants with data on Syphilis is 107, Haemoglobin at enrollment 424 and malaria is 302, this needs to be explained by the authors. Patients with malaria, low haemoglobin and syphilis are at risk for preterm birth, so missing data will influence results for the ZAPPS study.

This should be discussed by the authors as a study limitation if these figures are correct. These figures are correct. As we note above, since our study is integrated with routine care at the clinical sites, for several cases—particularly in the early conduct of the study—we did not repeat routine tests previously performed at government clinics and failed to document these results on our study forms. This is a limitation of the study that we have described in our Discussion (lines 295-299). We also note that it has been rectified in Phase 2 of the cohort (line 299).

This comment does not influence this paper as it for those recruited up to Sept 2017. Of note is that the protocol for ZAPPS has been amended in Feb 2018 to exclude women with HIV. So eventually will there be separate analysis for the women who are included in this manuscript vs those recruited later?

That is correct. Given the amended protocol has revised key inclusion / exclusion criteria, the next phase of the ZAPPS cohort has not been described in this baseline manuscript. We have added an explicit description of the two phases of the ZAPPS cohort in the Methods section (lines 102-104).

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