STUDY PROTOCOL

A Randomized controlled trial of the Effect of intraVenous iron on Anaemia in Malawian Pregnant women (REVAMP): Statistical analysis plan [version 2; peer review: 2 approved]

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First published: 07 Dec 2021, 5:174
https://doi.org/10.12688/gatesopenres.13457.1
Latest published: 14 Apr 2022, 5:174
https://doi.org/10.12688/gatesopenres.13457.2

Abstract

Background: Anaemia affects more than half of Africa’s pregnancies. Standard care, with oral iron tablets, often fails to achieve results, with compliance and gastrointestinal side-effects being a significant issue. In recent years, intravenous iron formulations have become safe, effective, and quick to administer, allowing the complete iron requirements of pregnancy to be provided in one 15-minute infusion. The Randomized controlled trial of the Effect of intravenous iron on Anaemia in Malawian Pregnant women (REVAMP) will evaluate whether a modern intravenous iron formulation, ferric carboxymaltose (FCM), given once during the second trimester is effective and safe in improving maternal and neonatal outcomes for treatment of moderate to severe anaemia in sub-Saharan Africa.

The objective was to publish the detailed statistical analysis plan for the REVAMP trial prior to unblinding the allocated treatments and
performing the analysis.

**Methods:** REVAMP is a multicentre, two-arm, open-label, parallel-group randomized control trial (RCT) in 862 pregnant women in their second trimester. The trial statistician developed the statistical analysis plan in consultation with the trial management team based on the protocol, data collection forms, and study outcomes available in the blinded study database.

**Results:** The detailed statistical analysis plan will support the statistical analyses and reporting of the REVAMP trial after unblinding the treatment allocations.

**Conclusions:** A statistical analysis plan allows for transparency as well as reproducibility of reporting and statistical analyses.

**Keywords**
anaemia, iron deficiency, intravenous iron, ferric carboxymaltose (FCM), pregnancy, randomized controlled trial, statistical analysis, Malawi
Amendments from Version 1

The revisions to version 2 of this manuscript are:
- Added a reference for the cut-off point for the primary outcome of maternal anaemia and clarified its definition across different timepoints
- Added secondary neonatal continuous outcomes for weight and length at 1 month postpartum to complement the infant growth z-scores
- Added that sample size calculations were performed using Stata software
- Clarified primary analyses will be undertaken on an intention-to-treat (ITT) basis and the definition of the ITT and per-protocol populations throughout
- Added a modified safety population definition excluding women who did not have a singleton pregnancy
- Clarified the model for the primary and secondary repeated timepoint binary outcomes, added the measure of effect for a modified Poisson regression model and added insight into why we selected a log-binomial model rather than a logistic regression model
- Clarified the description for the reporting and analysis of adverse events
- Clarified we will refer to the analysis as a modified ITT analysis if not all randomised participants will have data contributing to the analysis due to missing data
- Clarified about signing of the final version of SAP
- Revised the CONSORT flow chart footnotes to indicate those who were included in the ITT population

Version 1 of this manuscript is an extension of version 1 REVAMP SAP document dated October 2021. Version 2 of this manuscript is an extension of version 2 REVAMP SAP dated April 2022. The latter version incorporates the above listed changes to version 1 that were documented before unblinding of the study database and thus considered as planned analyses.

Any further responses from the reviewers can be found at the end of the article

Introduction

Approximately 36.5% of pregnant women globally are anaemic (World Health Organization, 23 April 2021), and iron-deficiency anaemia (IDA) is the cause of almost half of all anaemia during pregnancy1. In sub-Saharan Africa, 46% of all pregnant women are anaemic1. The adverse outcomes of anaemia during pregnancy extend to both the mother – including the life-threatening complication of postpartum haemorrhage – and the baby – including prematurity, low birth weight2, impaired development3, and increased mortality4,5. Thus, reducing the burden of anaemia in women is one of the key World Health Organization (WHO) 2025 global nutrition targets6.

Oral iron is the established approach for preventing and treating IDA in pregnancy and infancy7. However, oral iron may be poorly tolerated due to gastrointestinal adverse events8 and poorly adhered to over an entire course of treatment. This may result in suboptimal adherence to prevention programs in low- and middle-income countries8,9. Over the past two decades, parenteral (intravenous) iron therapies have dramatically advanced in terms of safety and convenience, providing an alternative to oral therapy. Modern parenteral iron formulations are commonly used in high-income countries to treat IDA during pregnancy10. Ferric carboxymaltose (FCM) is an established modern intravenous iron drug, which enables up to 1000 mg of elemental iron to be delivered in a single 15-minute infusion (NPS Medicinewise, 01 May 2021). FCM is approved for use in pregnancy after the first trimester11. The safety and convenience of FCM make this drug an exciting opportunity to treat anaemia in pregnancy in low-income countries. However, the evidence for the efficacy and safety of delivering FCM in pregnancy in low- to middle-income countries remains limited.

The Randomized controlled trial of the Effect of intraVenous iron on Anaemia in Malawian Pregnant women (REVAMP) is an open-label randomized controlled trial conducted in the Blantyre and Zomba districts of Malawi designed to determine the efficacy and safety of delivering FCM (compared with standard-of-care oral iron) in women with moderate or severe anaemia in the second trimester of pregnancy12 (ACTRN12618001268235). The primary outcome of the trial is maternal anaemia at 36 weeks’ gestation, and the key neonate outcome is birth weight. The trial recruited the first participant in November 2018 and completed follow-up to one-month postpartum in September 2021.

This paper describes the planned analysis for the REVAMP trial. This statistical analysis plan supersedes the plan provided in the trial registry and published protocol13. Finalization of the statistical analysis plan before study unblinding has been undertaken to ensure transparency in the methods used to analyze and report the data and ultimately create the evidence for the effects of intravenous iron supplementation on recovery from prenatal anaemia, haemoglobin, iron status, postpartum haemorrhage, and delivery outcomes.

Methods

The trial protocol is summarised elsewhere13.

Aims

The study’s main objective is to determine the efficacy and safety of a single intravenous iron administration during the second trimester of pregnancy – given as ferric carboxymaltose compared with routinely delivered oral iron – given as ferrous sulphate – in improving maternal (primarily anaemia) and neonatal (e.g., birth weight) outcomes.

Design

REVAMP is an open-label two-arm parallel-group randomized controlled trial in anaemic pregnant women (capillary haemoglobin <10 g/dL). Women were randomized to either IV ferric carboxymaltose 1000 mg (for women with weight >50 kg), or 20 mg/kg (for women with weight <50 kg) once during the second trimester; or oral iron 200 mg ferrous sulphate (approx. 65 mg elemental iron) twice daily for 90 days or the duration of pregnancy, whichever was shorter. Study visits occurred over pregnancy, at birth, and follow-up to one-month postpartum (Figure 1). The study received ethics approval from The College of Medicine Research Ethics Committee.
COMREC), Blantyre, Malawi (COMREC: P.02/18/2357) and the Walter and Eliza Hall Institute of Medical Research Human Ethics Committee, Melbourne, Australia (WEHI REC:18/02). It was prospectively registered at the Australian New Zealand Clinical Trials Registry (ACTRN12618001268235, registered on 27 July 2018).

**Setting**

The trial took place in southern Malawi at two sites: the coordinating site in Zomba Central Hospital, Zomba district, and a second site in Blantyre district at Limbe Health Centre. Both sites had all the resources required to recruit eligible participants, prepare, and administer the study drugs, monitor safety, treat adverse effects, and measure trial outcomes.

**Participants**

Women eligible for enrolment were in their second trimester (between 13–26 weeks of gestation) and presented with a capillary haemoglobin level below 10 g/dL, as measured by HemoCue® Hb 301 system. In addition, participants were eligible if they were negative for malaria (determined using a rapid diagnostic test (RDT)), planned to deliver at the health facility and were able to provide written informed consent (or have a legal guardian do so if <18 years old). Women with clinical symptoms of infection, any severe condition requiring hospitalization, a history of pre-eclampsia, or known hypersensitivity to the study drugs were not eligible for recruitment.

**Randomization and treatment allocation**

Women were randomly allocated to one of the two treatment arms with 1:1 allocation using a randomization schedule of randomly permuted blocks stratified by site to achieve balance between the arms within each site. The randomization list was computer-generated by an independent statistician and participants were randomly allocated using sealed, opaque envelopes. Although the trial is open-label, laboratory scientists measuring haemoglobin concentration, midwives collecting birth outcome data, and investigators and researchers in Australia (including data managers and statisticians in Melbourne) are blinded to the treatment allocation during the conduct of the trial until the database is locked and ready for unblinding.

**Outcome variables**

All efficacy and laboratory outcomes were measured at baseline, 28 days post randomization, 36 weeks’ gestation, delivery and one month postpartum for mothers and at delivery and one month postpartum for neonates. Data related to safety, including non-serious adverse events (AEs) and serious adverse events (SAEs), were collected across the total study period.
Maternal
The primary outcome of the study was maternal anaemia (defined as a venous haemoglobin concentration less than 11.0 g/dL) at 36 weeks’ gestation.

Secondary maternal outcomes included laboratory indices (haemoglobin and ferritin concentrations) and haematological and iron diagnoses (anaemia, moderate/severe anaemia, iron deficiency, iron deficiency anaemia) at 28 days post randomization, 36 weeks’ gestation, and one month postpartum. Haemoglobin concentration, anaemia and moderate/severe anaemia were also included at delivery. Using mother’s haemoglobin (g/dL) anaemia was defined as haemoglobin concentration less than 11.0 g/dL up to and including delivery and 12.0 g/dL postpartum, moderate/severe anaemia as haemoglobin concentration less than 10.0 g/dL and 11.0 g/dL respectively. Using mother’s ferritin (ug/L), and C-reactive protein (CRP, mg/L), iron-deficient was defined as ferritin <15 mg/L adjusted for inflammation (CRP >5 mg/L), and iron deficiency anaemia as iron deficient and anaemic.

Maternal safety outcomes included reported adverse events (including serious adverse events defined as any adverse event that resulted in death, were life threatening, required either inpatient hospitalization or prolongation of hospitalization, resulted in a persistent or significant disability/incapacity or resulted in a congenital anomaly/birth disorder), all-cause sick visits (including specifically, visits due to clinical malaria), hypophosphatemia (mild: 0.64< phosphate (PO4) <0.80 mmol/L, moderate: 0.32<PO4<0.64 mmol/L, severe: PO4 <0.32 mmol/L), inflammation (elevated C-reactive protein), and severe medical events (includes haemorrhage, need for transfusion, ICU admission, mortality). Adverse events were coded using version 5.0 of the Common Terminology Criteria for Adverse Events (CTCAE) (US Department of Health Human Services, 27 November 2020) and consisted of selecting an appropriate Preferred Term (PT) and System Organ Class (SOC) for each AE verbatim term. All AEs were coded on an ongoing basis by two physicians independent from each other and without knowledge of the treatment assignment (i.e., blinded). Any discrepancies regarding the selection of an appropriate PT or SOC will be resolved via discussion between the two physicians. All AE coding will be finalised before unblinding of the study database.

Additional data collected included maternal baseline characteristics: age, parity, gravidity, anthropometry, religion, education, marital status, income source, recent malaria status, and HIV positivity status.

Neonate
The key neonatal outcome was birth weight, measured in grams, within 24 hours of birth.

Secondary neonatal outcomes included birth length, gestation duration, and adverse birth outcomes (e.g., low birth weight, premature birth, small for gestation age, foetal loss – both individually and as a composite) within 24 hours of birth and growth (weight, length, weight-for-age z-score, length-for-age z-score, weight-for-length z-score) and laboratory indices (haemoglobin concentration) at one month of age. Using gestational duration (weeks), premature birth was defined as a neonate born before 37 completed weeks of gestation, foetal loss was defined as pregnancy loss before 28 completed weeks’ gestation or stillbirth as the birth of a child showing no signs of life after 28 weeks’ gestation. Low birth weight was defined as a neonate born with a birth weight <2500 g. Small for gestational age was derived using the neonate’s gestation duration and birth weight together with the sex of the child according to INTERGROWTH-21 standards. Growth outcomes (z-scores) were derived using the neonate’s length and weight together with age and sex of the neonate according to age and sex specific WHO international growth standards.

Neonate safety outcomes included (serious) adverse events, all cause sick visits (includes infection-related visits, diarrhoea-related visits, clinical malaria-specific visits).

Sample size
The sample size for the trial was to recruit 862 pregnant women (431 per arm) to have 80% power to detect that FCM will result in a 10% improvement in anaemia cure (a prevalence of 60% for standard oral therapy and 50% for FCM)\(^{12,17}\), allowing for a two-sided alpha of 5% and a 10% loss to follow-up. The sample size also has at least 80% power to detect an absolute difference between standard-of-care oral iron and FCM of 100 g in the neonatal outcome of birth weight, assuming a standard deviation of 450 g and a two-sided alpha of 5%, similar to the effect seen in a trial of Kenyan women receiving oral iron when compared with placebo\(^{9}\). No interim analyses to stop the trial early were planned, and no interim analysis was conducted. Sample size calculations were performed using Stata/SE (StataCorp. 2019. College Station, TX: StataCorp LLC).

Statistical analysis plan
The analysis will be conducted by statisticians at The Walter and Eliza Hall Institute of Medical Research (WEHI), Melbourne, Australia. After all study data are available and clean, a blinded data review meeting to review protocol violations and missing data will be held prior to database lock. The final statistical analysis plan will be signed off during this meeting. Analysis will be conducted using Stata/SE (StataCorp. 2019. College Station, TX: StataCorp LLC). The analysis of the primary outcome will be independently checked. Discrepancies will be discussed and resolved by consensus.

General principles
Maternal. Primary analyses will be undertaken on an intention-to-treat basis. The intention-to-treat population will consist of all mothers who were randomized and included in the analysis of all primary and secondary maternal outcomes according to the randomized allocation. The per-protocol population will consist of all mothers who were randomized, and without protocol violations. A protocol violation is defined as those who have withdrawn informed consent for the use of all their data or violating inclusion/exclusion criteria (e.g., twin pregnancy). These protocol violations are based
on pre-randomisation characteristics only and expected to be balanced between treatment groups. Mothers who are found to be non-adherent to treatment (e.g., refused or not provided treatment after randomisation) will be excluded from the per-protocol population analysis. Data on adherence to standard-of-care is restricted due to the suspension of home visits (Figure 1) to mitigate risks during the COVID-19 pandemic. The safety population will consist of all women who received at least one study treatment (either IV iron or standard-of-care, as treated) and included in the analyses of all safety maternal outcomes according to their actual treatment. A modified safety population was defined similar to the safety population and excluding women who had a singleton pregnancy protocol violation. Mothers who have withdrawn consent for use of all their data will be excluded from all analyses.

**Neonate.** Primary analyses will be undertaken on an intention-to-treat basis. The intention-to-treat population will consist of all live-born neonates (with the exception of the stillbirth outcome) of mothers who were randomized and included in the analysis of the key and secondary neonatal outcomes according to the mother’s randomized allocation. The per-protocol population will consist of all neonates born to mothers who were randomized, and without protocol violations. A protocol violation is defined as those who have withdrawn informed consent for the use of all their data or the mother being a protocol violation. Neonates of mothers who are found to be non-adherent to treatment (e.g., refused or not provided treatment after randomisation) will be excluded from the per-protocol population analysis. The safety population will consist of all neonates born to mothers who received at least one treatment (either IV iron or standard-of-care, as treated) and included in the analyses of all safety neonatal outcomes according to the actual treatment of the mother. A modified safety population was defined similar to the safety population and excluding neonates born to mothers who had a singleton pregnancy protocol violation. Neonates who have had consent withdrawn for use of all their data will be excluded from all analyses.

**Trial profile**
The flow of mothers and their neonates through the trial will be presented in a Consolidated Standards of Reporting Trials (CONSORT) diagram, reasons for exclusion will be reported (Figure 2).

**Baseline characteristics**
Maternal demographic and baseline variables will be summarised descriptively and presented by treatment group. No formal comparisons will be made between treatment arms. Characteristics will be summarised using frequencies and percentages (based on the non-missing sample size) for categorical variables, mean and standard deviation for continuous variables, or median and quartiles (25th and 75th percentile) for non-normally distributed continuous variables.

**Efficacy outcomes: analysis**

**Maternal.** Maternal anaemia (primary outcome) at 28 days post randomization, 36 weeks’ gestation (primary timepoint), delivery and 28 days postpartum will be analyzed using a log-binomial regression model, including mothers as a random intercept to account for multiple time points. The model will include a treatment and treatment by study visit interaction and adjust for the stratification variable used during the randomization (site). The model will include the standard-of-care oral iron group as the reference group. The treatment effect will be estimated from this model as the prevalence ratio of IV iron versus standard-of-care oral iron. In case of non-convergence, we will fit a modified Poisson regression model with robust error variance, including mothers as a random intercept to account for the multiple timepoints. The treatment effect will be estimated from this model as the prevalence ratio of IV iron versus standard-of-care oral iron. We have selected the prevalence ratio as an effect measure because its interpretation as the ratio change in prevalence is easier to understand than the interpretation of the odds ratio for clinical researchers.

Secondary repeated time-point binary outcomes (moderate/severe anaemia, iron deficiency, iron deficiency anaemia) will be analyzed similarly to anaemia at 28 days post randomization, 36 weeks’ gestation, delivery (for moderate/severe anaemia), and 28 days postpartum.

Secondary repeated time-point continuous outcomes (haemoglobin concentration and ferritin concentration) will be analyzed using a likelihood-based longitudinal data analysis model10. The model will assume a common baseline mean across the two treatment arms and an unstructured variance-covariance among the repeated measurements. The model will incorporate time point (study visit) as a categorical variable, treatment and treatment by study visit interaction and adjust for the stratification factor (site) as main effects. In case of non-convergence, we will consider alternative structures (first-order autoregressive, Toeplitz, compound symmetry). The treatment effect will be estimated from this model as the mean change from baseline to 28 days post randomization, 36 weeks’ gestation, delivery, and 28 days postpartum respectively between IV iron and standard-of-care oral iron. Ferritin (µg/L) will be log transformed before analysis, with the treatment effect expressed as a geometric mean ratio.

**Neonate.** Birth weight will be analyzed by fitting a linear regression model, adjusting for the stratification factor (site). The treatment effect will be estimated from this model as the absolute difference in mean birth weight between IV iron and standard-of-care oral iron.

Secondary single time point continuous outcomes (gestation duration and birth length within 24 hours of birth, and haemoglobin concentration, length-for-age z-score, weight-for-age z-score, and weight-for-length z-score at one month of age) will be analyzed similarly to birth weight. Appropriate transformations may be applied to the variables before fitting the model if considered skewed.

Secondary single time point binary outcomes (composite adverse birth outcome, low birth weight, foetal loss, premature birth, and small for gestational age) will be analyzed using a
Figure 2. The CONSORT flow chart.
log-binomial regression model, adjusting for the stratification factor (site). The treatment effect will be estimated from this model as the risk ratio of IV iron versus standard-of-care oral iron. In case of non-convergence, we will fit a modified Poisson regression model with robust error variance.

Safety outcomes: analysis

**Maternal.** The number and percentage of women who died, reported at least one serious adverse event (overall, and for each of the following time intervals: within 24 hours of randomization, within 14 days of randomization, antenatal and postpartum), reported at least one adverse event (overall, and for each of the following time intervals: within 24 hours of randomization, within 14 days of randomization, antenatal and postpartum), who had at least one severe medical event (composite, and its components haemorrhage, need for transfusion, ICU admission, or mortality), had at least one common AE (>5% in any group), had at least one AE by system organ class and had at least one AE by preferred term will be reported. Each of the above listed adverse event outcomes will be compared between treatment arms using a log-binomial regression model. In case of non-convergence for a safety outcome(s), a Poisson model with robust standard errors will be used to analyze the data. Prevalence ratios and 95% confidence intervals will be presented for IV iron versus standard-of-care oral iron. In addition, infusion-related adverse events will be reported separately for the IV iron group only.

The number and percentage of women with unplanned clinic visits (including all-cause sick visits and clinical malaria specific visits) and safety biomarkers (including inflammation and malaria RDT positive) will be reported and compared between treatments (by timepoint) using a log-binomial regression model. In case of non-convergence, a Poisson model with robust standard errors will be fitted to the data. The number and percentage of women with the safety biomarker hypophosphatemia will be reported and compared between treatments (by timepoint) using an ordered logistic regression model.

**Neonate.** The number and percentage of neonates who died, had at least one serious adverse event, had at least one adverse event, had at least one common AE (>5% in any group), had at least one AE by system organ class and had at least one AE by preferred term will be reported and compared between treatments using a log-binomial regression model. In case of non-convergence, a Poisson model with robust standard errors will be used.

The number and percentage of neonates with unplanned clinic visits (including all-cause sick visits, diarrhoea related visits, respiratory related visits and clinical malaria specific visits) will be reported and compared between treatments using a log-binomial regression model. As for other binary outcomes, a Poisson model with robust standard errors will be fitted if there are non-convergence issues when fitting a log-binomial regression model.

Reporting and methods for missing data

To describe the missing data, the frequency and percentage of study participants with missing data at baseline, 28 days post randomization, 36 weeks’ gestation, delivery and one month postpartum will be summarised for anaemia (mothers) and birth weight (neonates) by treatment group. In addition, baseline and demographic characteristics will be summarised by those with baseline only, incomplete data at any visit, and complete data at all visits for anaemia (mothers) to explore the missing data assumption(s) and identify any study variables not included in the target analyses that are potentially associated with missing/not missing of these study variables (known as auxiliary variables). As a rule of thumb, if the proportion of missing data is below approximately 5%, those values will be considered negligible in the case of maternal anaemia or live-born neonates with missing birth weight\(^2\), in which case we will refer to the analysis as a modified intention-to-treat analysis.

For dealing with missing data in the analyses of primary and secondary outcomes, the primary analysis will be an available case analysis performed for repeated time point outcomes (e.g., anaemia) and a complete case analysis for single time point outcomes (e.g., birth weight).

**Maternal.** As the primary strategy to handle missing data, the analysis of maternal anaemia (repeated assessments) will use a likelihood-based approach. This approach relies on the underlying assumption that the probability of missing outcome data is not related to the missing data after conditioning on observed data in the model (Missing at Random [MAR]).

If the missing data is not negligible, additional analysis will be performed whereby missing maternal haemoglobin data will be multiply imputed using chained equations, separately by treatment group. The imputation model will include site, parity, gestational age at baseline, body mass index (BMI), inflammation status at baseline, iron deficient status at baseline, HIV positive status at baseline, and re-screened post positive malaria RDT status at baseline.

In addition, auxiliary variables identified during the blinded data review meeting may be included. Maternal haemoglobin will be imputed using a linear regression model. The missing outcome data at 28 days post randomization, 36 weeks’ gestation, delivery and one month postpartum will be imputed using the “just another variable” approach (also known as imputing in wide format), which requires a separate imputation model for imputing the variable at each assessment time\(^3\). The number of imputed data sets will be greater than or equal to the percentage of missing data in the available case analyses. Using these imputed data sets, an analysis based on a pattern-mixture model\(^4\) consisting of applying a delta-adjustment to the imputed values by treatment group will be conducted. Within the standard-of-care group participants with missing data will be assumed having both a poorer and better response than those with observed data while no a priori difference is anticipated in the mean response for the IV iron group. Differences in baseline participant characteristics between those with and without data will inform delta-values to explore in both treatment groups. After deriving maternal anaemia from the imputed haemoglobin values, the imputed data sets will be analyzed using a log-binomial regression model. The estimates from the analyses of the imputed data
sets will be combined to obtain a pooled common estimate and corresponding confidence interval for the effect of the iron intervention on maternal anaemia using Rubin’s rules. The delta-adjustment method within the multiple imputation framework assumes a Missing Not At Random (MNAR) assumption for the outcome.

**Neonate.** The analysis of birth weight will use a complete-case analysis among those live-born. This approach relies on the underlying assumption that the probability of missing outcome data is not related to the observed or missing data (Missing Completely at Random [MCAR]).

If the missing data is not negligible, additional analysis will be performed whereby missing birth weight data will be multiply imputed, separately by treatment group. The imputation model will include site, sex, gestational age at baseline and maternal BMI. In addition, auxiliary variables identified during the blinded data review meeting may be included. Birth weight will be imputed using a linear regression model. The number of imputed data sets will be greater than or equal to the percentage of missing data in the complete case analyses. The estimates from the analyses of the imputed data sets will be combined to obtain a pooled common estimate and corresponding confidence interval for the effect of the iron intervention on birth weight using Rubin’s rules. This approach relies on the MAR assumption for the outcome, birth weight.

**Additional analyses**

**Maternal.** In addition to the analysis model for all maternal efficacy outcomes adjusted for site as a main effect, additional analyses will be performed for these outcomes:

1. Analyses consisting of models adjusted for auxiliary variables:
   a. Adding to the model adjusted for site, the main effect of parity (primiparous vs. multiparous), gestational age at baseline (continuous), and BMI at baseline (continuous).
   b. Adding to the model adjusted for site, parity, gestational age at baseline and BMI, the main effects of inflammation status at baseline, iron-deficient status at baseline, haemoglobin at baseline (continuous), HIV positive status at baseline, and re-screened post-previous positive malaria RDT status.
   c. Adding to the model adjusted for site, the main effect of variables in demographic and/or baseline characteristics demonstrating imbalance between treatment arms after unblinding.

2. Analysis of the model adjusted for site for the per-protocol population.

3. Analysis of the model adjusted for site for the per-protocol population adjusted for baseline characteristics demonstrating imbalance between treatment arms.

Furthermore, we will report the number-needed-to-treat (NNT) and 95% confidence interval for maternal anaemia at 36 weeks’ gestation.

**Neonate.** In addition to the analysis model for all neonate efficacy outcomes adjusted for site as a main effect, additional analysis will be performed for these outcomes:

1. Analyses consisting of models adjusted for auxiliary variables:
   a. Adding to the model adjusted for site, the main effect of sex of the infant (female or male), gestational age at baseline (continuous), and maternal BMI at baseline (continuous).
   b. Adding to the model adjusted for site, sex, gestational age at baseline and maternal BMI the main effect of maternal haemoglobin at baseline (continuous).
   c. Adding to the model adjusted for site, the main effect of variables in demographic and/or baseline characteristics demonstrating imbalance between treatment arms after unblinding.

2. Analysis of the model adjusted for site for the per-protocol population.

3. Analysis of the model adjusted for site for the per-protocol population adjusted for baseline characteristics considered not balanced between the arms for the per-protocol population.

Furthermore, we will report the NNT and 95% confidence interval for low birth weight.

**Multiple testing**

No adjustment for multiplicity is planned for the primary maternal outcome (anaemia at 36 weeks’ gestation) and key neonate outcome (birth weight). We will test the primary null hypothesis of no difference between IV iron and standard-of-care oral iron at a two-sided 5% level of significance. Estimates and two-sided confidence intervals will be presented, along with multiplicity unadjusted P-Values.

The Holm procedure is used to ensure control of the Type I error rate for secondary maternal outcomes at 36 weeks’ gestation (haemoglobin concentration, moderate/severe anaemia, ferritin, iron deficiency, and iron deficiency anaemia) and neonate outcomes (gestation duration, birth length, composite adverse birth outcome within 24 hours of birth and infant growth at 1-month postpartum) separately. We will present multiplicity unadjusted P-values along with the estimate and 95% confidence intervals and footnote the comparisons meeting the statistical significance threshold according to the Holm procedure.

No multiplicity adjustment is planned for other secondary outcomes at 28 days post randomization, delivery, and 28 days postpartum. We will present the estimate and two-sided 95% confidence interval and no P-Values will be presented. We will present the multiplicity unadjusted P-Values for the safety outcomes; no multiple testing adjustment is planned.

**Subgroup analyses**

Exploratory subgroup analyses will be performed for the outcomes of maternal anaemia at 36 weeks’ gestation, haemoglobin concentration at 36 weeks’ gestation, birth weight, low
birth weight, gestation duration, and premature birth. The following subgroups will be explored: parity (primiparous vs. multiparous), baseline HIV status (positive vs negative), baseline severe anaemia status (yes vs no severe anaemia), baseline iron-deficient status (yes vs no ID), baseline iron-deficient anaemia status (yes vs no IDA), baseline inflammation status (yes vs no elevated CRP), re-screened after positive malaria RDT at pre-screening (yes vs no) and site (Blantyre, Zomba). In addition, subgroup (main effect) and the subgroup-by-treatment interactions term will be added to the unadjusted model to evaluate whether the treatment effect (IV iron versus standard-of-care) differs between subgroup categories. No multiplicity adjustments are planned for the subgroup analyses due to their exploratory nature. Results of the subgroup analyses (effect estimate and 95% Confidence Interval) will be displayed using Forest plots.

**Trial status**

This statistical analysis plan is an extension of the REVAMP protocol and documents version 1 dated October 26, 2021. Any changes to this version between publishing and unblinding will be tracked and still considered as planned analyses. The final statistical analysis plan will be approved during the blinded data review and signed before breaking the allocation code, after which any changes will be considered post-hoc.

**Discussion**

Antenatal anaemia remains a significant public health concern in low-to-middle income countries. Although oral iron supplementation remains a cheap formulation, suboptimal adherence and common limiting gastrointestinal adverse effects from the drugs may limit effectiveness. If our data demonstrate a benefit from intravenous iron on maternal outcomes and potentially also on critical neonatal outcomes such as birth weight, the findings will provide evidence for the beginning of a clinical rationale for developing strategies for implementing this intervention in practice. Thus, results from this trial could ultimately transform the way anaemia is treated in low-income settings and have long term benefits for maternal and child health, ultimately resulting in benefits for maternal and child survival.

**Data availability**

No data are associated with this article.

**Acknowledgements**

The authors thank the participants and their families involved in the study.

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Open Peer Review

Current Peer Review Status: ✔ ✔

Version 1

Reviewer Report 04 January 2022

https://doi.org/10.21956/gatesopenres.14714.r31529

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Tapan Rai

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This is a well-written statistical analysis plan which is written as concisely as possible. Although the publication of statistical analysis plans for clinical trials has become more common over the past few years, it is not yet standard practice and the art of writing a plan for publication as a journal article is still evolving. This article does a very good job of this and may be used as a guide by other research teams. Having said that, I believe that there may be some places where the authors could clarify a few points - especially since other teams are likely to look at this plan as a model in the future:

1. The Intention-to-Treat (ITT) and per-protocol cohorts could be better defined. I would specifically seek clarification of the following points:
   1. The authors state: "The per-protocol population will consist of all mothers who were randomized, and without protocol violations. A protocol violation is defined as no informed consent ...". This seems to make it clear that participants who did not provide informed consent will be excluded from the per-protocol cohort. However, it also seems to suggest that they will be included in the ITT cohort. This is not usually the case, since the trial should not generally be collecting data from participants who have not or are unable to provide informed consent - unless informed consent is provided by proxy (e.g. by next-of-kin). If the authors did not intend to suggest that participants without informed consent are to be included in the ITT cohort, please reword or restate the definition of a protocol violation so as to remove any doubt about this. Alternatively, can the authors, please, clarify whether:
      1. The trial collected data from participants without informed consent?
      2. If so, was consent obtained from next-of-kin?
      3. Are participants who did not provide informed consent to be included in the ITT cohort?
   2. If a participant refused treatment after being randomised or were otherwise not
given the allocated treatment in the expected timeframe, they would normally be analysed in the group to which they are randomised in an ITT analysis. However, it is not clear how these participants would be analysed in the per-protocol analysis - would they be deleted from the analysis or analysed with the standard care group (e.g. if they were randomised to the FCM group but received standard care)?

2. The proposed log-binomial models (with possible use of a Poisson model with robust errors, if the log-binomial model fails to converge) are appropriate in this situation. However, since statistical analysis plans are often read by clinical researchers (who are not all statisticians), please write 1-2 sentences explaining your rationale for selecting this approach rather than a logistic regression model which is more common (and more clinical researchers are likely to be familiar with). My hope is that this will provide some guidance/insight on this to other researchers reading this statistical analysis plan.

3. The statistical analysis plan for secondary repeated timepoint continuous outcomes, mentions the use of the interaction effect between treatment group and timepoint (p. 7, Paragraph 3). This is standard practice to assess between-group differences at specific timepoints in a longitudinal model. I expect that you are using a similar approach (interaction between treatment group and timepoint) for the primary and secondary repeated timepoint binary outcomes as well. However, you have not mentioned this in the text. For consistency and clarity, please state this clearly when outlining the analysis for these binary outcomes as well (alternatively, if you are not using some other approach that does not require the interaction effect in these models, please specify this and elaborate on this so that it is clear to the reader).

4. The description for the reporting and analysis of adverse events needs some clarification. There are several adverse event outcomes mentioned (for maternal adverse events). These include the number and percentage of women who:
   1. died,
   2. reported at least one serious adverse event (including within 24 hours of randomization, within 14 days of randomization, antenatal and postpartum),
   3. reported at least one adverse event (including within 24 hours of randomization, within 14 days of randomization, antenatal and postpartum),
   4. at least one severe medical event (composite, and its components haemorrhage, need for transfusion, ICU admission, or mortality),
   5. had at least one common AE ,
   6. had at least one AE by system organ class and had at least one AE by preferred term will be reported.

This suggests that these will be reported separately for each stated timepoint (e.g. at least one AE within 24 hours, at least one AE in the period 24 hours-14 days post randomization etc.) and by organ system etc. Please clarify if this is the case, and, if this is the case, please clarify what response variable will be used for the log-binomial model associated with adverse events - is it one model for each AE outcome mentioned above (which is not likely to converge if the number of events in each category is small) or is there some designated response variable to be used?

Overall, it may be easier to clarify how the adverse events would be reported by providing sample tables of the reporting of descriptive statistics as an appendix. Providing such sample tables would
also help make clearer how some of the other descriptive statistics will be reported.

**Is the rationale for, and objectives of, the study clearly described?**
Yes

**Is the study design appropriate for the research question?**
Yes

**Are sufficient details of the methods provided to allow replication by others?**
Yes

**Are the datasets clearly presented in a useable and accessible format?**
Not applicable

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

**Reviewer Expertise:** Biostatistics

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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Author Response 07 Apr 2022

**Rebecca Harding**

Thank you for your review and feedback comments

Responses to your minor comments:

1. Thank you for pointing this out. We did not intend to suggest that participants without informed consent are to be included in the ITT population. The definition has been reworded in the general principles section:
   “A protocol violation is defined as those who have **withdrawn informed consent for the use of all their data** or violating inclusion/exclusion criteria (e.g., twin pregnancy).”
   To clarify: The trial did not collect data from participants without informed consent. Participants who did not provide informed consent will not be included in the ITT population.
2. Thank you for pointing this out. The following text has been added in the general principles section.
   Maternal - Mothers who are found to be non-adherent to treatment (e.g., refused or not provided treatment after randomisation) will be excluded from the per-protocol population analysis.
   Neonate - Neonates of mothers who are found to be non-adherent to treatment (e.g., refused or not provided treatment after randomisation) will be excluded from the per-protocol population analysis.

2.
The following text has been added to the efficacy outcomes analyses section. We have selected the prevalence ratio as an effect measure because its interpretation as the ratio change in prevalence is easier to understand than the interpretation of the odds ratio for clinical researchers.

3. Thank you for pointing this out, yes, we are using a similar approach of a treatment and treatment by study visit interaction for the primary repeated timepoint binary outcomes as well. The following bold text has been added to the efficacy outcomes: analyses section: Maternal anaemia (primary outcome) at 28 days post randomization, 36 weeks' gestation (primary timepoint), delivery and 28 days postpartum will be analyzed using a log-binomial regression model, including mothers as a random intercept to account for multiple time points. The model will include a treatment and treatment by study visit interaction and adjust for the stratification variable used during the randomization (site).

4. The analysis will be performed by adverse event outcome with the response variable being the binary adverse event outcome (e.g. died: yes/no, had at least one common AE: yes/no). We have changed and added the following bold text to the safety outcomes: analyses section:

The number and percentage of women who died, reported at least one serious adverse event (overall, and for each of the following time intervals: within 24 hours of randomization, within 14 days of randomization, antenatal and postpartum), reported at least one adverse event (overall, and for each of the following time intervals: within 24 hours of randomization, within 14 days of randomization, antenatal and postpartum), who had at least one severe medical event (composite, and its components haemorrhage, need for transfusion, ICU admission, or mortality), had at least one common AE (>5% in any group), had at least one AE by system organ class and had at least one AE by preferred term will be reported. Each of the above listed adverse event outcomes will be compared between treatment arms using a log-binomial regression model. In case of non-convergence for a safety outcome(s), a Poisson model with robust standard errors will be used to analyze the data. Prevalence ratios and 95% confidence intervals will be presented for IV iron versus standard-of-care oral iron. In addition, infusion-related adverse events will be reported separately for the IV iron group only.

Thank you for your suggestion of providing sample tables in an appendix. We hope to have satisfactorily clarified our approach with regards to the adverse event safety outcomes in text.

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

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The Statistical analysis plan has been well constructed with excellent statistical methodology. There are a few areas that need to be addressed to improve the plan as follows:

Minor comments:

1. The plan clearly defines the Intention to treat and per protocol populations, however, there is no commitment in the plan as to which of the two approaches will form the primary strategy for the primary outcome. Is it the ITT or the PP? This should be clarified. Simply defining intention to treat or per-protocol populations without further specification is not sufficient.

2. The plan is clear that for the primary outcome, a log-binomial regression model will be used or in case of non-convergence, a modified Poisson regression model with robust error variance will be fitted. However, the measure of effect has not been clearly stated. For a non-technical audience, it is important to state the measure of effect like how the authors have stated for neonatal birth weight.

3. In the Trial status section, authors need to state about signing of the final version of SAP by the investigators.

4. The authors state that if the proportion of missing data will be less than 5%, available case and complete case analysis approaches will be used. While I agree with this rule of thumb, I am wondering how authors will handle the intention to treat analyses if there will be less than 5% missing outcome data. Will they analyse fewer number of participants than randomized? If so what will be the implications of the ITT principle?

5. Will “mixed effects” be considered in any of the specified statistical models? The authors should state in the SAP if these are anticipated.

6. The CONSORT flow chart has been very well constructed and is clear. However, there is no indication on which levels constitute ITT analyses and which ones constitute PP analyses.

7. In the sample size section, the software that was used for sample size calculations should be stated.
8. On page 4, the authors state that “The primary outcome of the study was maternal anaemia (defined as a venous haemoglobin concentration less than 11.0g/dL) at 36 weeks’ gestation.” It would be good to insert the reference for the cut-off point.

Is the rationale for, and objectives of, the study clearly described?
Yes

Is the study design appropriate for the research question?
Yes

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

Are the datasets clearly presented in a useable and accessible format?
Not applicable

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

**Reviewer Expertise:** Statistics, epidemiology, clinical trials

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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Author Response 07 Apr 2022

**Rebecca Harding**

Thank you for your review and feedback comments

Responses to your minor comments:

1. Thank you for pointing this out. This has been clarified in the general principles section. The following bold text has been added:
   *Maternal. Primary analyses will be undertaken on an intention-to-treat basis. The intention-to-treat population will consist of all mothers who were randomized…*

   *Neonate. Primary analyses will be undertaken on an intention-to-treat basis. The intention-to-treat population will consist of all live-born neonates (with the exception of the stillbirth outcome) of mothers who were randomized and included in the analysis of the key and secondary neonatal outcomes according to the mother’s randomized allocation.*

2. This point has been addressed in the efficacy outcomes: analysis section. The following bold text has been added:
   “In case of non-convergence, we will fit a modified Poisson regression model with robust error variance, including mothers as a random intercept to account for the multiple timepoints. **The treatment effect will be estimated from this model as the prevalence ratio of IV iron versus standard-of-care oral iron.**”
3. This point has been addressed and the following bold text has been added in the trial status section:
   “The final statistical analysis plan will be approved during the blinded data review and signed before breaking the allocation code, after which any changes will be considered post-hoc.”

4. If the analysis of the primary maternal endpoint (anaemia at 36 weeks’ gestation) or the key neonate endpoint (birthweight) result in exclusion of participants due to missing all outcome data, then we will refer to this analysis as a modified intention-to-treat analysis. The following bold text has been added to the reporting and methods for missing data section:
   As a rule of thumb, if the proportion of missing data is below approximately 5%, those values will be considered negligible in the case of maternal anaemia or live-born neonates with missing birth weight [19], in which case we will refer to the analysis as a modified intention-to-treat analysis.

5. The specified log-binomial regression model for repeated timepoint binary outcomes and the specified likelihood-based longitudinal data analysis model for repeated time point continuous outcomes contain both fixed and random effects.

6. A footnote has been added to the CONSORT flow chart to indicate those who were included in the Intention-to-Treat (ITT) population.

7. The following text has been added in the sample size section:
   “Sample size calculations were performed using Stata/SE (StataCorp. 2019. College Station, TX: StataCorp LLC).”

8. The reference for the cut-off point has been added.
   World Health Organization. Haemoglobin concentrations for the diagnosis of anaemia and assessment of severity. Vitamin and Mineral Nutrition Information System. Geneva, World Health Organization, 2011 (WHO/NMH/NHD/MNM/11.1) (http://www.who.int/vmnis/indicators/haemoglobin.pdf, accessed 17 Feb 2022).

Competing Interests: No competing interests were disclosed.