Intermittent Preventive Therapy Post-Discharge: an innovative approach in the prevention of rebound severe malaria anaemia and mortality in young children

Submission date	Recruitment status No longer recruiting	Prospectively registered	
18/05/2006		☐ Protocol	
Registration date	Overall study status	Statistical analysis plan	
19/06/2006	Completed	[X] Results	
Last Edited	Condition category	[] Individual participant data	
10/04//01/	INFECTIONS AND INFESTATIONS		

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Dr Feiko ter Kuile

Contact details

Liverpool School of Tropical Medicine Pembroke Place Liverpool United Kingdom L3 5QA +44 (0)151 7053287 terkuile@liverpool.ac.uk

Additional identifiers

Protocol serial number N/A

Study information

Scientific Title

Intermittent Preventive Therapy Post-Discharge: an innovative approach in the prevention of rebound severe malaria anaemia and mortality in young children - a randomised double-blind placebo controlled multicentre study

Acronym

IPTpd

Study objectives

To compare the efficacy of a single treatment course with lumefantrine-artemether (Coartem®) at discharge to three treatment courses with Coartem® given at discharge, 1 and 2 months (intermittent preventive therapy post-discharge [IPTpd]), to standard antimalarial therapy of oral sulfadoxine-pyrimethamine (SP) in Malawi, in the post-discharge management of children, aged 4-59 months, who have recovered from severe malarial anaemia by assessing mean haemoglobin concentration, and the incidence of rebound severe anaemia, clinical malaria and death by 3 and 6 months.

As of 22/04/2010 this record was updated; all changes can be found in the relevant fields with the above update date. At this time, the anticipated end date of this trial was also updated; the initial anticipated end date at the time of registration was 01/12/2008.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved by College of Medicine Research and Ethics Committee on 25/02/05, reference number: P.03/04/287 and by Liverpool Research and Ethics Committee on 09/02/05, reference number: 05.01

Study design

Randomised, double-blind, placebo-controlled, multicentre study

Primary study design

Interventional

Study type(s)

Prevention

Health condition(s) or problem(s) studied

Severe malarial anaemia

Interventions

Patients are randomised into one of the following groups:

Group A - lumefantrine-artemether, single 3-day course at enrolment

Group B - lumefantrine-artemether, three 3-day courses (at enrolment, at 1 month, and at 2 months)

Group C - sulfadoxine-pyrimethamine (SP), single dose at enrolment (added 22/04/2010: group C dropped out following amendments to protocol)

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

lumefantrine-artemether (Coartem®), sulfadoxine-pyrimethamine

Primary outcome(s)

Current information as of 22/04/2010:

The incidence of rebound severe anaemia (Hb less than 5 g/L), severe malaria (hospital admissions requiring quinine) or death (a composite endpoint) between 1 and 6 months after enrolment.

Initial information at time of registration:

Mean haemoglobin at three months

Key secondary outcome(s))

Current information as of 22/04/2010:

- 1. The incidence of sick-child's clinic visits due to clinical malaria by 3 and 6 months
- 2. The incidence of all-cause sick-child's clinic visits by 3 and 6 months
- 3. The incidence of all cause re-hospitalisation between 1 3 and 1 6 months after enrolment
- 4. The incidence of the three individual components of the composite endpoint (severe anaemia, severe malaria, death) between 1 3 and 1 6 months after enrolment
- 5. Mean haemoglobin at 6 months
- 6. Incidence of adverse events by 3 and 6 months
- 7. Mean corrected heart rate (QTc) prolongation by 3 days

Initial information at time of registration:

- 1. The incidence of sick-child's clinic visits due to clinical malaria by 3 and 6 months
- 2. The incidence of rebound severe anaemia (Hb <5 g/l)
- 3. The incidence of death by 3 and 6 months
- 4. Mean haemoglobin at 6 months
- 5. Incidence of adverse events by 3 and 6 months
- 6. Mean corrected heart rate (QTc) prolongation by 3 days

Completion date

01/12/2010

Eligibility

Key inclusion criteria

- 1. Haemoglobin <5.0 g/dl or packed cell volume (PCV) <15% on admission to the hospital
- 2. Plasmodium falciparum malaria (any documented parasitaemia) at the time of admission to the hospital or within 24 hours prior to admission
- 3. Aged between 4 months (inclusive) and 59 months (inclusive) at the time of randomization
- 4. Bodyweight >5 kg at the time of randomization
- 5. Subject completed blood transfusion(s) in accordance with routine hospital practice
- 6. Subject completed intravenous (IV) quinine in accordance with routine hospital practice
- 7. Able to feed (for breastfed children) or eat (for older children)
- 8. Able to sit unaided
- 9. Provision of informed consent by parent or guardian

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Child

Lower age limit

4 months

Upper age limit

59 months

Sex

All

Key exclusion criteria

- 1. Recognised, specific other cause of severe anaemia at the time of admission to the hospital (e.
- g. trauma, haematological malignancy, known bleeding disorder, known sickle cell disease)
- 2. Previous enrolment in the present study
- 3. Severe anaemia (haemoglobin <5.0 g/dl) at the time of randomization
- 4. Known hypersensitivity to any of the study drugs
- 5. Documented intake of Coartem® (≥4 doses) or SP within 1 week prior to admission
- 6. Child resides outside of catchment area during the course of the study (6 months)
- 7. Known need at the time of randomization for concomitant prohibited medication during the 2 months randomized treatment period
- 8. Ongoing participation into another clinical trial involving ongoing or scheduled treatment with medicinal products during the course of the study (6 months)
- 9. Known need, or scheduled surgery during the course of the study (6 months)
- 10. Suspected non-compliance with the follow-up schedule

Date of first enrolment

22/05/2006

Date of final enrolment

01/12/2010

Locations

Countries of recruitment

United Kingdom

England

Malawi

Study participating centre **Liverpool School of Tropical Medicine** Liverpool **United Kingdom** L3 5QA

Sponsor information

Organisation

Liverpool School of Tropical Medicine (UK)

ROR

https://ror.org/03svjbs84

Funder(s)

Funder type

Research organisation

Funder Name

The Netherlands-African partnership for capacity development and clinical interventions against poverty-related diseases (NACCAP) (Netherlands) (ref: W 07.05.202.00)

Funder Name

UBS Optimus Foundation (Switzerland)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	Details	Date created Date added	Peer reviewed	Patient-facing?
Results article	results	01/03/2012	Yes	No

Participant information sheet Participant information sheet 11/11/2025 No

Yes