Impact of early administration of artesunate suppository on childhood severe malaria in Tanzania

Submission date 01/02/2006	Recruitment status No longer recruiting	[] Pr [] Pr
Registration date 01/02/2006	Overall study status Completed	[_] St [X] R
Last Edited 23/02/2009	Condition category Infections and Infestations	[_] In(

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s) Scientific

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers 980482

Prospectively registered

] Protocol

Statistical analysis plan

X] Results

[_] Individual participant data

Study information

Scientific Title

Study objectives

The objective has been to establish whether, in patients with acute malaria who cannot take medication by mouth, rectal artesunate plus referral differs from rectal placebo plus referral in terms of death or permanent disability.

Ethics approval required Old ethics approval format

Ethics approval(s) Ethics approval received on the 1st January 1999.

Study design Randomised controlled trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Not specified

Study type(s) Treatment

Participant information sheet

Health condition(s) or problem(s) studied Malaria

Interventions

The sample size determination in the protocol specified that a total of 10,000 non per os patients would need to be randomised in order to detect a reduction of mortality from 5% to 3%.

Individual patients will be randomised to receive either AS suppository (intervention group) or placebo (comparator group). Patients in both groups will then be referred immediately to the nearest hospital/health centre.

Intervention Type

Drug

Phase Not Specified

Drug/device/biological/vaccine name(s)

Artesunate (AS)

Primary outcome measure

1. Number of relevant deaths in the intervention and control arm assessed 7 - 30 days after enrolment (relevant defined as malaria positive patients in whom the death was probably /definitely preventable by the intervention)

2. Number of individuals with serious neurological disability in the intervention and control arms assessed at 7 - 30 days following enrolment in the study. Neurological disability defined as the development of new problems with feeding, walking, talking, sitting, sight, hearing, playing, balance and behaviour

Secondary outcome measures

1. Number of deaths in the intervention and control arm assessed 7 - 30 days following enrolment in the study

2. Number of cases of neurological disability in the intervention and control arms assessed at 7 - 30 days following enrolment in the study

3. Number of cases of neurological disability in malaria smear positive patients in the intervention and control arms assessed at 7 - 30 days following enrolment in the study 4. Number of cases of neurological disability in children in the intervention and control arms assessed at 7 - 30 days following enrolment in the study

5. Number of cases of neurological disability in pregnant women in the intervention and control arms assessed at 7 - 30 days following enrolment in the study

6. Number of deaths and neurological sequelae in the intervention and control arm in malaria smear positive patients who survived at least 8 hours but died before 7 days after enrolment in the study

Overall study start date

01/01/1999

Completion date

01/01/2001

Eligibility

Key inclusion criteria

1. Non per os children presenting to a peripheral health unit or traditional healer with clinicallly suspected P. falciparum malaria

2. Children (apparent age 6 months to 71 months) with suspected malaria who are too ill to take drugs by mouth

3. Individual informed consent - consent must have been given (generally by thumb-print) by the patient or guardian of the sick person

4. Community informed consent - at the start of the study in that area, community consent to the project would have been obtained

Participant type(s) Patient

Age group Child

Lower age limit

6 Months

Upper age limit

71 Months

Sex Not Specified

Target number of participants

In this trial, it is not the number of patients recruited but the number of deaths that determine the statistical power of such a trial.

Key exclusion criteria Ability to take an oral medication.

Date of first enrolment 01/01/1999

Date of final enrolment 01/01/2001

Locations

Countries of recruitment Switzerland

Tanzania

Study participating centre 20, Avenue Appia Geneva-27 Switzerland CH 1211

Sponsor information

Organisation

UNICEF/UNDP/World Bank/WHO Special Programme for Research and Training in Tropical Diseases (TDR)

Sponsor details

20, Avenue Appia Geneva-27 Switzerland CH 1211

Sponsor type Research organisation

Website http://www.who.int/

ROR https://ror.org/01f80g185

Funder(s)

Funder type Research organisation

Funder Name

United Nations Children's Fund (UNICEF)/United Nations Development Programme (UNDP) /World Bank/World Health Organization (WHO) - Special Programme for Research and Training in Tropical Diseases (TDR)

Funder Name European Commission (Belgium)

Alternative Name(s)

European Union, Comisión Europea, Europäische Kommission, EU-Kommissionen, Euroopa Komisjoni, Ευρωπαϊκής Επιτροπής, Εвροπεйската комисия, Evropské komise, Commission européenne, Choimisiúin Eorpaigh, Europskoj komisiji, Commissione europea, La Commissione europea, Eiropas Komisiju, Europos Komisijos, Európai Bizottságról, Europese Commissie, Komisja Europejska, Comissão Europeia, Comisia Europeană, Európskej komisii, Evropski komisiji, Euroopan komission, Europeiska kommissionen, EC, EU

Funding Body Type

Government organisation

Funding Body Subtype National government

Location

Funder Name

WHO Global Malaria Programme

Funder Name US Agency for International Development (USAID) (USA)

Funder Name Irish Aid (Ireland)

Funder Name Karolinska Institutet (Sweden)

Alternative Name(s) Karolinska Institute, KI

Funding Body Type Government organisation

Funding Body Subtype Local government

Location Sweden

Funder Name Sall Family Foundation (USA)

Alternative Name(s)

Funding Body Type Private sector organisation

Funding Body Subtype Trusts, charities, foundations (both public and private)

Location United States of America

Funder Name University of Oxford Clinical Trial Service Unit (UK)

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	
Results article	

Details Date created results 14/02/2009 Date added Peer reviewed?

Yes

Patient-facing?

No