

Comparison of how well allopathic and herbal medicine work for the treatment of *Entamoeba histolytica*

Submission date 19/07/2012	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
Registration date 30/07/2012	Overall study status Completed	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
Last Edited 13/10/2016	Condition category Infections and Infestations	<input type="checkbox"/> Individual participant data <input type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

Amoebiasis is an infection of the gut caused by a parasite called *Entamoeba histolytica*, which is found in most tropical areas. Infections occur in both the intestine and (in people with symptoms) in the tissue of the intestine and/or liver. As a result, two different classes of drugs are needed to treat the infection, one for each location, such as metronidazole and diloxanide furoate. It is claimed that a herbal medicine (Endemali) treats both types of infection with very few side effects. The aim of this study is to find out which of the two treatments is better with respect to effectiveness and side effects in order to find an alternative treatment for *E. histolytica*, which is developing growing resistance against the traditional medicine.

Who can participate?

Patients aged 5 to 60 with amoebiasis

What does the study involve?

Participants are randomly allocated to one of two treatment groups. One group was treated with a combination of metronidazole and diloxanide furoate three times a day for 5 days. The other group was treated with Endemali four times a day for 10 days. During the course of treatment, participants are checked for the development of side effects or any other complications. At the end of the study participants are checked for the presence or absence of *E. histolytica*.

What are the possible benefits and risk of the participating?

This study could help patients in the future by providing an option of a drug which is more effective and has fewer side effects. Participants are provided with treatment and their investigations free of charge. Those for whom the treatment did not work are further investigated and provided with other treatment options without charge. There is a chance of developing side effects and the drugs may not work.

Where is study run from?

Hamdard University (Pakistan)

When is the study starting and how long is it expected to run for?
January to December 2008

Who is funding the study?
Hamdard University (Pakistan)

Who is the main contact?
1. Prof. Dr Muhammad Irfanullah Siddiqui (irfan7255@yahoo.com)
2. Prof. Dr Usman Ghani (ugk_2005@yahoo.com)

Contact information

Type(s)
Scientific

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers
001

Study information

Scientific Title
A randomized double blind study to find out whether there is any difference in the efficacy and side effects of Endemali® and a combination of diloxanide furoate and metronidazole

Study objectives
There are two hypothesis simultaneously tested:
1. It is hypothesized that there is some difference between Endamali (a herbal medicine) and a combination of metronidazole and diloxanide furoate with respect to efficacy and side effects. (Two-tailed)

Null hypothesis: There is no difference in the efficacy of Endamali and a combination of metronidazole and diloxanide furoate with respect to efficacy.

2. It is hypothesized that there is some difference between Endamali (a herbal medicine) and a combination of metronidazole and diloxanide furoate with respect to side effects. (Two-tailed)

Null hypothesis: There is no difference in the efficacy of Endamali and a combination of metronidazole and diloxanide furoate with respect to side effects.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics Committee of Hamdard University, 16/01/2007, ref: CMHCMD/001/2007

Study design

Double-blind randomized parallel bi-centric study

Primary study design

Interventional

Secondary study design

Randomised parallel trial

Study setting(s)

Hospital

Study type(s)

Treatment

Participant information sheet

Not available in web format, please use the contact details to request a patient information sheet

Health condition(s) or problem(s) studied

Amoebiasis

Interventions

To determine the sample size, we used $\alpha < 0.05$ and power of study as 90%. The effect size considered to be clinically significant, was a 0.15 difference between two groups based on a pilot study. Hypothesis test for two population proportion with two sided test was calculated with the help of WHO Manual. A total of 101 sample size was calculated for each group. The total 202 patients were divided into 21 blocks. The block technique used has already been described in the application sent to you, in which we used the block of 10 in order to have almost equal number of participants in both group. Out of the 10 sheet written for each block, five were marked "treatment group 1" (TR1) and rest as "treatment group 2" (TR2). Each eligible participant was invited to pick blindly one sheet out of 10 available. Once the five patients of one treatment group were complete, rest of the patients were allocated to the other group e.g. in one draw first patient picked TR1, second also picked TR1, 3rd picked TR2, 4th picked TR1 5th picked TR2, 6th picked TR2, 7th picked TR2, 8th picked TR1, 9th picked TR1 and then 10th was automatically allocated to TR2. Once a sheet was picked, after noting the group allocation, it

was put back in the drawer to make it 10 again so that every patient has an equal probability of being allocated to any of the group.

Treatment group 1: combination of metronidazole + diloxanide furoate available in Pakistan with brand name "Entamizole DS". This tablet contains metronidazole 400mg + diloxanide furoate 500mg. Entamizole DS was given 3 times a day for 5 days.

Treatment group 2: Endemali®, a herbal product, available in 4gm sachet, containing Boswellia glabra 270.9 mg, Kaolinum ponderosum 255 mg, Ocimum pilosum 580 mg, Pistacia terbinthus 116.1 mg, Plantago ispagula 812.7 mg, Vateria indica 232.2mg sweetening agent q.s. Endemali® was given 4 times a day for 10 days.

Intervention Type

Drug

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

Diloxanide furoate, metronidazole

Primary outcome measure

Efficacy at baseline and 5 days after stopping the treatment

Secondary outcome measures

1. Side effects
2. Association of age with efficacy
3. Association of sex with efficacy
4. Association of age with side effects
5. Association of sex with side effects

Measured at baseline and immediately after completion of course of treatment.

Overall study start date

01/01/2008

Completion date

31/12/2008

Eligibility

Key inclusion criteria

1. All the confirmed cases of amoebiasis and between the ages of 5 and 60 years were invited to be included in the study
2. Willing to be assigned to any of the two treatment groups

Participant type(s)

Patient

Age group

Adult

Sex

Both

Target number of participants

202

Key exclusion criteria

All those who had multiple problems

Date of first enrolment

01/01/2008

Date of final enrolment

31/12/2008

Locations**Countries of recruitment**

Pakistan

Study participating centre

House No A-4 ST-33

Karachi

Pakistan

75850

Sponsor information**Organisation**

Hamdard University (Pakistan)

Sponsor details

Madinat Al Hikmah

Muhammad Bin Qasim Avenue

Near Bund Murad

Gadap town

Karchi

Pakistan

74600

Sponsor type

University/education

Website

<http://www.hamdard.edu.pk/>

ROR

<https://ror.org/01zrv0z61>

Funder(s)

Funder type

University/education

Funder Name

Hamdard University (Pakistan)

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