

# Fever in Subsaharan Africa

<b>Submission date</b> 02/08/2019	<b>Recruitment status</b> No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 14/10/2019	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 15/04/2025	<b>Condition category</b> Signs and Symptoms	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Fever is a common symptom that is defined as a higher than normal body temperature. It occurs most often in response to an infection. There are many infections that can cause fever. These infections can be due to many different types of germs (bacteria, viruses, parasites and fungi). ALERRT (The "African coalition for Epidemic Research, Response and Training") is a group of health institutions that have come together to conduct research that aims to improve care and treatment for people who have an infectious disease in Africa.

### Who can participate?

The FISSA study is about adults and children above two months presenting with a febrile illness and without hospitalization in the 14 last days.

### What does the study involve?

The FISSA study is an observational study which aims to better understand the main causes of fever, by describing symptoms and signs of your health problem including its severity, the care received and the outcome of this care. The researcher will not influence the patient's care, but simply records what is happening. Medical staff who attend to participants will care for them in the same way as usual. A better understanding of these features will lead to development of new treatments and improve care.

### What are the possible benefits and risks of participating?

The care that the participant will receive is exactly the same he/she would receive outside of the study context. It will entail no additional cost to and no blood test or additional examination will be required. The only additional study-specific procedure will be phone calls or on site visits, for which the participant will receive a mobile phone credits/units or a small cash for transportation compensation.

### Where is the study run from?

The Study will take place in 16 care centers in sub Saharan Africa (8 in Western, 4 in Central and 4 in Eastern Africa). Recruitment will take place in 11 countries and 8 sites.

### When is the study starting and how long is it expected to run for?

The first recruitment is planned for November 25, 2019 and the study is scheduled to last 12 months. Each participant will be followed up for 21 days.

Who is funding the study?

The FISSA Study is funded by European Union through a European and Developing Countries Clinical Trials Partnership (EDCTP) Program.

Who is the main contact?

Anani Badje

Coordinator of de Work Package 1 of ALERRT Network

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## Contact information

### Type(s)

Public

### Contact name

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Scientific

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### Contact details

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Non applicable

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## Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

### ClinicalTrials.gov (NCT)

Nil known

**Protocol serial number**

Nil known

## **Study information**

**Scientific Title**

Clinical pattern, severity, management and outcome of acute febrile illness in sub-Saharan Africa

**Acronym**

FISSA

**Study objectives**

The field of infectious diseases is diverse and fast-moving. Researchers should be prepared to tackle diseases which pose major threats, such as HIV/AIDS, tuberculosis or malaria, but also new emerging diseases before they become major public health threats, such as viral hemorrhagic fevers. African clinical trials network for infectious diseases should be adaptable and have both a generic approach to clinical research and a specific approach depending on the disease. This implies building: (i) networks of clinical centres across as many countries as possible where professionals have been trained in clinical trials and are able to implement standard procedures, (ii) networks of regional and international institutions that bring them in contact with the best specialists in specific fields. To consolidate the setting of this network, to consolidate the setting of this network, the FISSA study is being implemented with the following objectives to document the clinical patterns, severity, management, and outcomes of febrile illnesses in clinical centers across West, Central and East Africa.

**Ethics approval required**

Old ethics approval format

**Ethics approval(s)**

Approved 26/09/2019, Comite National d'Ethique des Sciences de la Vie et de la Sante (CNESVS s/c de l'institut pasteur, 01 BP 490 Abidjan 01, Ivory Coast; +225 41 40 05 55; cnesvscotedivoire@gmail.com), ref: 106-19

**Study design**

Multi-center prospective observational cohort study

**Primary study design**

Observational

**Study type(s)**

Other

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

Febrile illnesses

**Interventions**

After giving informed consent to participate in the study, each participant will be included and followed for 21 days. Contacts are planned for days 7, 14 and 21 after inclusion. These contacts will be made by telephone or through visits as far as possible. However, unscheduled calls or visits may be made at any time if necessary.

## **Intervention Type**

Other

## **Primary outcome(s)**

Clinical status is measured using a three-category ordinal scale (resolved, not resolved, dead) at day 7.

## **Key secondary outcome(s)**

1. Clinical status is measured using a three-category ordinal scale (resolved, not resolved, dead) at day 14 and day 21.
2. Illness severity:
  - 2.1. In adults:
    - 2.1.1. Ambulatory Simplified Acute Physiologic Score (ASAPS) and modified ASAPS (using ACVPU instead of Glasgow) at days 0, 7, 14 and 21.
    - 2.1.2. Quick Sepsis Related Organ Failure Assessment (qSOFA) at days 0, 7, 14 and 21.
  - 2.2. In children <5 years is measured using the ALgorithm for the MANAgement of Childhood illness (ALMANACH) criteria at days 0, 7, 14 and 21.
3. The proportion of subjects admitted to hospital is measured at day 21.
4. Length of stay in hospital (Initial and/or secondary hospitalization) is measured at day 21.
5. Number of secondary visits is measured at day 21.
6. Number and type of laboratory and radiology tests prescribed/actually performed measured at day 21.
7. Number and type of medications and supportive care prescribed/actually given measured at day 21.
8. Working diagnoses at admission and final diagnoses measured at day 21.
9. Patient satisfaction is measured using a questionnaire at day 21.

## **Completion date**

25/11/2020

## **Eligibility**

### **Key inclusion criteria**

1. Ongoing objectively determined fever, as defined by axillary Temperature  $>37.5^{\circ}\text{C}$ , or tympanic (or oral or rectal) temperature  $>38^{\circ}\text{C}$ .
2. Contactable by the study team at days 7, 14 and 21.
3. Informed consent to participate signed by the patient (adults) or a legally acceptable representative (children or patients with impaired consciousness) with assent from the concerned, where applicable and possible

## **Participant type(s)**

Patient

## **Healthy volunteers allowed**

No

## **Age group**

Mixed

## **Sex**

All

**Total final enrolment**

8867

**Key exclusion criteria**

1. Children under 2 months
2. Hospitalization in the last 14 days

**Date of first enrolment**

25/11/2019

**Date of final enrolment**

10/12/2021

## **Locations**

**Countries of recruitment**

Cameroon

Central African Republic

Côte d'Ivoire

Gambia

Ghana

Guinea

Madagascar

Nigeria

Senegal

Tanzania

Uganda

**Study participating centre**

Children's Hospital of Diamniadio

Diamniadio

Diamniadio

Senegal

Non applicable

**Study participating centre**  
**CHU de Fann/Albert Royer**  
Dakar  
Dakar  
Senegal  
Non applicable

**Study participating centre**  
**Centre de Santé amélioré de Maferinyah**  
Maferinyah  
Maferinyah  
Guinea  
Non applicable

**Study participating centre**  
**CHR d'Ayamé**  
Ayamé  
Ayamé  
Côte d'Ivoire  
Non applicable

**Study participating centre**  
**Urgences du CHU de Treichville**  
Treichville  
Côte d'Ivoire  
Non applicable

**Study participating centre**  
**St. Francis Xavier Hospital**  
Assin Foso  
Assin Foso  
Ghana  
Non applicable

**Study participating centre**  
**Federal Medical Centre of Owo**  
Owo  
Owo  
Nigeria  
Non applicable

**Study participating centre**  
**Clinical Service Department Fajara**  
Fajara  
Fajara  
Gambia  
Non applicable

**Study participating centre**  
**Les Promoteurs de la Bonne Santé**  
Yaoundé  
Yaoundé  
Cameroon  
Non applicable

**Study participating centre**  
**Obala District Hospital**  
Obala  
Obala  
Cameroon  
Non applicable

**Study participating centre**  
**Hôpital de District de Sibut**  
Sibut  
Sibut  
Central African Republic  
Non applicable

**Study participating centre**  
**Boda District Hospital**  
Boda  
Boda  
Central African Republic  
Non applicable

**Study participating centre**  
**CHU de Befelatanana**  
Befelatanana  
Befelatanana

Madagascar  
Non applicable

**Study participating centre**

**HDSS Moramanga**

Moramanga

Moramanga

Madagascar

Non applicable

**Study participating centre**

**Korogwe District Hospital**

Korogwe

Korogwe

Tanzania

Non applicable

**Study participating centre**

**Uganda Virus Research Institute Clinic**

Entebbe

Entebbe

Uganda

Non applicable

## **Sponsor information**

**Organisation**

EDCTP

**ROR**

<https://ror.org/031jv9v19>

## **Funder(s)**

**Funder type**

Research organisation

**Funder Name**



## European and Developing Countries Clinical Trials Partnership

### Alternative Name(s)

Le partenariat Europe-Pays en développement pour les essais cliniques, A Parceria entre a Europa e os Países em Desenvolvimento para a Realização de Ensaaios Clínicos, The European & Developing Countries Clinical Trials Partnership, European and Developing Countries Clinical Trials, EDCTP

### Funding Body Type

Private sector organisation

### Funding Body Subtype

International organizations

### Location

Netherlands

## Results and Publications

### Individual participant data (IPD) sharing plan

Researchers can request ALERRT for access to the anonymized data for well-defined research or secondary analyses via a controlled access procedure. A Data Access Committee (DAC) will be set up. This Committee will review the request and provide timely an answer to the Requestor. In case of positive evaluations data will be shared by a data sharing agreement between the Platform Host and Requestor. Data will be shared from the Platform Host to Requestor by means of secure encrypted data transfer.

### IPD sharing plan summary

Available on request

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Participant information sheet</a>	Participant information sheet	11/11/2025	11/11/2025	No	Yes