Expanded safety and acceptability study of 6% cellulose sulphate

Submission date 19/03/2004	Recruitment status No longer recruiting	Prospectively registered
		☐ Protocol
Registration date	Overall study status	Statistical analysis plan
01/04/2004	Completed	[X] Results
Last Edited	Condition category	Individual participant data
10/11/2022	Urological and Genital Diseases	

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Dr Isaac Malonza

Contact details

World Health Organization 20 Avenue Appia Geneva-27 Switzerland CH-1211

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malonzai@who.int

Additional identifiers

Protocol serial number

WHO/HRP ID: A15006

Study information

Scientific Title

Expanded safety and acceptability study of 6% cellulose sulphate

Study objectives

Local tolerance and acceptability of cellulose sulphate (CS) gel applied vaginally twice daily for seven days.

Ethics approval required

Old ethics approval format

Ethics approval(s)

The study protocol was approved by scientific and ethics review committees at each implementing centre and the World Health Organization.

Study design

A phase I randomised, closed label, comparative study

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Sexually transmitted infection (STI) prevention

Interventions

- 1. Test groups using Cellulose Sulphate (CS) gel with or without concurrent intercourse
- 2. Control groups using K-Y jelly with or without concurrent intercourse

Total duration of involvement in the study, including screening, admission, completion and follow up, is approximately one month per subject.

Intervention Type

Drug

Phase

Phase I

Drug/device/biological/vaccine name(s)

Cellulose sulphate (CS) gel

Primary outcome(s)

The number of women who experienced any signs and/or symptoms of genital irritation as reported by volunteers at any time during follow-up or as determined by naked eye examination of the genitalia, on colposcopy or microbiologic tests.

Key secondary outcome(s))

Adverse events

Completion date

01/07/2003

Eligibility

Key inclusion criteria

- 1. Healthy, sexually abstinent and sexually active women
- 2. Considered to be at low risk for human immunodeficiency virus (HIV)
- 3. Recruited from family planning clinics and local communities in Kampala (Uganda), Mumbai (India), and Sagamu (Nigeria)
- 4. Aged between 18 and 50 years
- 5. Had regular menstrual cycles, or were on injectable contraceptives and amenorrhoeic for at least 6 months
- 6. Were not at risk for pregnancy (because of tubal ligation, steroidal contraceptives, or abstinence)
- 7. Willing to adhere to the study protocol

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

Female

Total final enrolment

180

Key exclusion criteria

- 1. Known to have an allergy to any component of CS gel or K-Y Jelly or condoms (cohort II only)
- 2. Currently pregnant or within 2 months from last pregnancy outcome
- 3. Known to abuse drug or alcohol
- 4. Had evidence of an infection with Trichomonas vaginalis, Candidiasis or bacterial vaginosis which did not resolve with treatment or if they had gonorrhoea or a chlamydial infection
- 5. Had a history of herpes or condylomata within the past 6 months
- 6. Had non-iatrogenic abnormal colposcopy findings involving deep disruption of the genital epithelium at enrolment

Date of first enrolment

01/12/2001

Date of final enrolment

01/07/2003

Locations

Countries of recruitment

India

Nigeria

Switzerland

Uganda

Study participating centre World Health Organization Geneva-27 Switzerland CH-1211

Sponsor information

Organisation

UNDP/UNFPA/WHO/World Bank - Special Programme of Research, Development and Research Training in Human Reproduction

ROR

https://ror.org/01f80g185

Funder(s)

Funder type

Research organisation

Funder Name

United Nations Development Programme (UNDP)/United Nations Population Fund (UNFPA) /World Health Organization (WHO)/World Bank - Special Programme of Research, Development and Research Training in Human Reproduction (HRP)

Results and Publications

Individual participant data (IPD) sharing plan

Not provided at time of registration

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output typeDetailsDate createdDate addedPeer reviewed?Patient-facing?Results article02/12/2005YesNo