Safety and efficacy of a cream on the wound healing and scar of cesarean section

Submission date 08/05/2024	Recruitment status No longer recruiting	 Prospectively registered Protocol
Registration date 10/05/2024	Overall study status Completed	 [] Statistical analysis plan [X] Results
Last Edited 27/01/2025	Condition category Pregnancy and Childbirth	Individual participant data

Plain English summary of protocol

Background and study aims

One of the most popular surgical procedures is the cesarean section (CS), and one of the most prevalent side effects of this procedure is wound and scar issues. The aim of this prospective, short-term, placebo-controlled study is to evaluate the safety and effectiveness of a silicone-based medical device (Lecoxen cream, Ekuberg Pharma, Italy) in healing wounds and scars derived from cesarean sections in nulliparous women. Methods:

Who can participate? Women, ranging in age from 18 to 45 years, who underwent CS

What does the study involve?

Participants are divided in two groups of treatment (Lecoxen cream or Vaseline). The quality of scar will be examined using the Vancouver scar scale after 28 days, while wound healing will be evaluated using the redness, edema, ecchymosis, discharge, and approximation (REEDA) scale on the 14th day following CS.

What are the possible benefits and risks of participating? Possible benefits: reduciton of healing time of wound and amelioration of scar tissue Possible risk: allergies to any of the ingredients contained in the proposed treatment

Where is the study run from? Department of Obstetrics and Gynecology "Veris delli Ponti Hospital" (Italy)

When is the study starting and how long is it expected to run for? June 2023 to December 2023

Who is funding the study? Investigator initiated and funded

Who is the main contact? Prof. Andrea Tinelli, andreatinelli@gmail.com

Contact information

Type(s) Scientific

Contact name Dr Davide Carati

ORCID ID http://orcid.org/0000-0003-2360-3456

Contact details Via castrignano Martano Italy 73025 +39 3294513080 davide.carati@ekubergpharma.com

Type(s) Public, Principal Investigator

Contact name Prof Andrea Tinelli

ORCID ID http://orcid.org/0000-0001-8426-8490

Contact details Direttore U.O.C. Ginecologia e Ostetricia, Ospedale Veris delli Ponti Scorrano Italy 73020 +39 3392074078 Ginecologia.poscorrano@asl.lecce.it

Additional identifiers

EudraCT/CTIS number Nil known

IRAS number

ClinicalTrials.gov number Nil known

Secondary identifying numbers 75839/2023

Study information

Scientific Title

Safety and efficacy of Lecoxen cream on the wound healing and scar of cesarean section: a prospective observational clinical trial

Study objectives

Lecoxen cream is more effective than pure Vaseline in reducing time of healing of wound resulting from cesarean section and improve quality of resulting scar.

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 06/06/2023, Ethical committee of Lecce (Via Miglietta 5, Lecce, 73100, Italy; +39 836 420273; cometico@asl.lecce.it), ref: 75839/2023

Study design Single center interventional randomized controlled trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Hospital

Study type(s) Safety, Efficacy

Participant information sheet

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

Health condition(s) or problem(s) studied

Treatment of wound derived after cesarean section

Interventions

All pregnant women submitted to a Pfannenstiel skin incision will be included in the study. After surgery, skin will be closured by agrafes (Agrafes Appose, Medtronic Italy, Milano, Italy) and will be divided into two 71 groups: 37 mothers were allocated to the treatment group with Lecoxen cream, 37 to the placebo group treated with 72 Vaseline (Vaseline F.U., Marco Viti, Italy). Participants were assigned in a 1:1 ratio with sequential assignment, depending on time and date of enrolment.

After the stitches' removal, normally on the seventh post-operative day, patients will be instructed to apply respective treatments from the first day to the twenty-eighth day, once a day. Patients will be asked not to use any other medicament for wound healing.

A board-certified gynecologist, all doctors working in the department where the study will be

performed, will supervised the screening, the enrollment of study participants from a panel of candidates, and will follow the patients during the study.

The state of wound healing will be evaluated at baseline and after 14 days (T1) of the treatment using Redness, Edema, Ecchymosis, Discharge, and Approximation (REEDA) scale. The degree of quality of the resulting scar will be finally evaluated after 28 days (T2), using Vancouver Scar Scale (VSS).

Intervention Type

Device

Pharmaceutical study type(s)

Dose response

Phase Not Specified

Drug/device/biological/vaccine name(s)

Lecoxen cream

Primary outcome measure

1. The quality of scar was examined using the VSS scale at baseline and after 28 days

2. The wound healing was evaluated using REEDA scale at baseline and after 14 days

Secondary outcome measures

Adverse events measured using patient records up to the end of the study

Overall study start date

01/06/2023

Completion date

31/12/2023

Eligibility

Key inclusion criteria Healthy women aged 18 - 45 years scheduled for CS

Participant type(s) Patient

Age group Adult

Lower age limit 18 Years

Upper age limit 45 Years

Female

Target number of participants

74

Total final enrolment

74

Key exclusion criteria

- 1. Smoking
- 2. At risk for premature birth
- 3. Obesity
- 4. Hypertension
- 5. Diabetes and infections
- 6. Use systemic and topical corticosteroids
- 7. Presence of previous surgical scarring near the incision site
- 8. Autoimmune diseases
- 9. Skin problems
- 10. Known hypersensitivity to at least one of the components of the formulation Lecoxen cream or Vaseline

Date of first enrolment

07/06/2023

Date of final enrolment 01/12/2023

Locations

Countries of recruitment Italy

Study participating centre Department of Gynecology and Obstetrics of the 'Veris Delli Ponti' Hospital Via Giuseppina Delli Ponti Scorrano Italy 73020

Sponsor information

Organisation Department of Obstetrics and Gynecology "Veris delli Ponti Hospital"

Sponsor details

Via Giuseppina Delli Ponti Scorrano Italy 73020 +39 836420564 ginecologia.poscorrano@asl.lecce.it

Sponsor type Hospital/treatment centre

Funder(s)

Funder type Other

Funder Name Investigator initiated and funded

Results and Publications

Publication and dissemination plan

Planned pubblication in a peer-reviewed journal

Intention to publish date 01/06/2024

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be available upon request from Andrea Tinelli andreatinelli@gmail.com

IPD sharing plan summary Available on request

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<u>Results article</u>		22/07/2024	27/01/2025	Yes	No