A single-centre study of the clinical efficacy of Eakin cohesive paste

Submission date	Recruitment status	[X] Prospectively registered
24/01/2005	No longer recruiting	☐ Protocol
Registration date	Overall study status	Statistical analysis plan
16/03/2005	Completed	☐ Results
Last Edited	Condition category	☐ Individual participant data
04/02/2014	Injury, Occupational Diseases, Poisoning	Record updated in last year

Plain English summary of protocol

Not provided at time of registration

Study website

http://www.eakin.co.uk

Contact information

Type(s)

Scientific

Contact name

Ms Marie McGrogan

Contact details

Royal Group of Hospitals Trust Grosvenor Road Belfast United Kingdom BT12 6BX

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers

TGE/P01

Study information

Scientific Title

Study objectives

Eakin cohesive paste is safe for use on patients with high output wounds and stomas.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Favourable ethical opinion received 16th September 2005.

Study design

Randomised controlled trial

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Not specified

Study type(s)

Not Specified

Participant information sheet

Health condition(s) or problem(s) studied

Stoma, ostomy, wound and fistula

Interventions

The trial is a randomised cross over trial. Patients will be assigned a patient identity number and will then be randomly

assigned either to begin with their normal pouching regime or their normal pouching regime and Eakin cohesive paste. At the half way point the treatment will be switched.

Intervention Type

Other

Phase

Not Specified

Primary outcome measure

Confirmation of the safety and efficacy of Eakin cohesive paste

Secondary outcome measures

Determination of whether Eakin cohesive paste increases the pouch weartime on high output wounds.

Overall study start date

01/06/2005

Completion date

01/10/2005

Eligibility

Key inclusion criteria

Patients presenting wounds, stomas and fistulae with greater than 200 ml exudate per day.

Participant type(s)

Patient

Age group

Adult

Sex

Both

Target number of participants

40

Key exclusion criteria

Patients will be excluded from the study under the following circumstances:

- 1. Where informed consent is withheld
- 2. Where the patient is unable to give informed consent due to legal incompetence
- 3. Where, in the physician's opinion, inclusion in the trial is not advised
- 4. Where the patient is presenting a critical wound, or is in an emergency situation
- 5. Where the patient is in the intensive care unit
- 6. Where there is an open wound in the peristomal region (for stoma patients only)
- 7. Where the patient is currently participating in another clinical trial
- 8. Where the patient has a known sensitivity to the product or any of its ingredients
- 8. Where the patient currently uses stoma paste as part of their regular pouching regime

Date of first enrolment

01/06/2005

Date of final enrolment

01/10/2005

Locations

Countries of recruitment

Northern Ireland

United Kingdom

Study participating centre Royal Group of Hospitals Trust Belfast United Kingdom BT12 6BX

Sponsor information

Organisation

T G Eakin Limited (UK)

Sponsor details

15 Ballystockart Road Comber United Kingdom BT23 5QY

Sponsor type

Not defined

Website

http://www.eakin.co.uk

Funder(s)

Funder type

Industry

Funder Name

T G Eakin Limited

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 summaryNot provided at time of registration