

An observer blind, parallel group, randomised multicentre study to compare the safety and efficacy of a new formulation of topical clindamycin phosphate in patients with mild to moderate acne

Submission date

09/09/2005

Recruitment status

No longer recruiting

Registration date

06/10/2005

Overall study status

Completed

Last Edited

17/09/2009

Condition category

Skin and Connective Tissue Diseases

☐ Prospectively registered

☐ Protocol

☐ Statistical analysis plan

☒ Results

☐ Individual participant data

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Prof WJ Cunliffe

Contact details

Department of Dermatology

Leeds General Infirmary

Leeds

United Kingdom

LS1 3EX

Additional identifiers

Protocol serial number

ZCG/4/C

Study information

Scientific Title

Study objectives

To ascertain the efficacy of a new formulation of topical clindamycin gel.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Not provided at time of registration

Primary study design

Interventional

Study design

Multicentre randomised active controlled parallel group trial

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Mild to moderate acne vulgaris.

Interventions

Zindaclin® 1% Gel (twice a day [bd] & once a day [od]) and Dalacin-T® Topical Lotion (bd).

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

Zindaclin® 1% Gel and Dalacin-T® Topical Lotion

Primary outcome(s)

The change in facial inflammatory lesion count from baseline visit to the end of treatment (week 16).

Key secondary outcome(s)

1. The change in acne grade (The Leeds Revised Acne Grading System, 1998) from the baseline visit to the end of treatment
2. The change in number of total comedones, open comedones, closed comedones, pustules
3. Mean change in total lesion count
4. Patients global assessment
5. Investigators global assessment

Completion date

29/09/2000

Eligibility

Key inclusion criteria

Consenting patients of either sex, between 12 and 40 years of age, with mild to moderate inflammatory acne with a grade ranging from 2.0 to 7.0 (The Leeds Revised Acne Grading System, 1998).

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Other

Sex

All

Key exclusion criteria

1. Patients with significant nodulocystic acne
2. Use of systemic topical antibiotics within 4 weeks prior to the start of treatment
3. Concomitant medication which may interfere with study evaluation

Date of first enrolment

19/07/1999

Date of final enrolment

29/09/2000

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

Department of Dermatology

Leeds

United Kingdom

LS1 3EX

Sponsor information

Organisation

ProStrakan Pharmaceuticals (UK)

ROR

<https://ror.org/017hh7b56>

Funder(s)**Funder type**

Industry

Funder Name

ProStrakan Pharmaceuticals (UK)

Results and Publications**Individual participant data (IPD) sharing plan****IPD sharing plan summary**

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/09/2005		Yes	No