

Study of COLAL-PRED® in the treatment of moderate acute ulcerative colitis

Submission date 10/03/2006	Recruitment status No longer recruiting	☐ Prospectively registered ☐ Protocol
Registration date 29/03/2006	Overall study status Completed	☐ Statistical analysis plan ☐ Results
Last Edited 31/01/2019	Condition category Digestive System	☐ Individual participant data ☐ Record updated in last year

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

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Additional identifiers

ClinicalTrials.gov (NCT)
NCT00299013

Protocol serial number
ATL2502/020/CL

Study information

Scientific Title
Study of COLAL-PRED® in the treatment of moderate acute ulcerative colitis

Study objectives

The aim of this study is to investigate whether COLAL-PRED® is non-inferior in terms of efficacy and superior in terms of safety to that of conventional prednisolone in the treatment of moderate acute ulcerative colitis.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Southampton and South West Hampshire Research Ethics Committees (A), 11/10/2005, ref: 05/Q1702/128

Primary study design

Interventional

Study design

Randomised double-blind double-dummy active-comparator parallel-group trial

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Moderate acute ulcerative colitis

Interventions

Patients are randomised to receive one of the following interventions:

1. Capsules containing prednisolone metasulfobenzoate sodium in a colonic delivery system
2. Prednisolone tablets
3. Matching placebo capsules and tablets

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

COLAL-PRED® containing prednisolone metasulfobenzoate sodium

Primary outcome(s)

1. Responder analysis for reduction in disease activity index (DAI) score
2. Responder analysis for safety based on morning cortisol levels

Key secondary outcome(s)

Treatment responder analysis of patients who are both efficacy and safety responders

Completion date

31/03/2007

Eligibility

Key inclusion criteria

1. Endoscopically confirmed diagnosis of ulcerative colitis
2. Score of 6-10 on the disease activity index (DAI)
3. Moderate to severe mucosal appearance

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

All

Key exclusion criteria

1. Previous colonic surgery
2. Other treatments for ulcerative colitis that have not been stabilised
3. Clinically significant diabetes
4. Heart failure
5. Unstable angina
6. Cirrhosis
7. Renal failure
8. History of tuberculosis

Date of first enrolment

07/03/2006

Date of final enrolment

31/03/2007

Locations**Countries of recruitment**

United Kingdom

England

Australia

Belgium

Czech Republic

Denmark

France

Germany

Hungary

Israel

Italy

Poland

Russian Federation

South Africa

Spain

Sweden

Study participating centre

University Hospital

Nottingham

United Kingdom

NG7 2UH

Sponsor information

Organisation

Alizyme (UK)

Funder(s)

Funder type

Industry

Funder Name

Alizyme (UK)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration