A multicentre, double-blind, randomised, parallel group, placebo-controlled trial of LAX-101 (ethyl eicosapentaenoate [EPA]) in patients with a new or recurrent episode of depression

Submission date 03/02/2003	Recruitment status No longer recruiting	 Prospectively registered Protocol
Registration date 03/02/2003	Overall study status Completed	 Statistical analysis plan Results
Last Edited 11/10/2016	Condition category Mental and Behavioural Disorders	 Individual participant data Record updated in last year

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Dr Harald Murck

Contact details

Laxdale Ltd Kings Park House Laurelhill Business Park Polmaise Road Stirling United Kingdom FK7 9JQ +44 (0)1786 476001 hmurck@laxdale.co.uk

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers LA01.01.0008A

Study information

Scientific Title

A multicentre, double-blind, randomised, parallel group, placebo-controlled trial of LAX-101 (ethyl eicosapentaenoate [EPA]) in patients with a new or recurrent episode of depression

Study objectives Not provided at time of registration

Ethics approval required Old ethics approval format

Ethics approval(s) Not provided at time of registration

Study design Randomised controlled trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Not specified

Study type(s) Treatment

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

Health condition(s) or problem(s) studied Depression

Interventions

Four groups randomly allocated to placebo, 0.5 g, 1 g or 2 g/day ethyl EPA on a double-blind basis.

Intervention Type

Other

Phase

Not Specified

Primary outcome measure Not provided at time of registration

Secondary outcome measures Not provided at time of registration

Overall study start date 01/01/2003

Completion date 01/01/2004

Eligibility

Key inclusion criteria

 Written informed consent
 Diagnosis of major depressive disorder (Diagnostic and Statistical Manual of Mental Disorders, Fourth edition [DSM-IV])
 Score of between and including 16 and 25 on the Hamilton Depression Rating Scale (HDRS)
 No treatment with any antidepressant medication (including St John's Wort) in the last 12 weeks from the date of Visit 0 (screen)
 Male or female of any race aged 18-75

Participant type(s) Patient

Age group Adult

Lower age limit 18 Years

Upper age limit 75 Years

Sex Both

Target number of participants Not provided at time of registration

Key exclusion criteria Not provided at time of registration

Date of first enrolment 01/01/2003

Date of final enrolment

01/01/2004

Locations

Countries of recruitment Scotland

United Kingdom

Study participating centre Laxdale Ltd Stirling United Kingdom FK7 9JQ

Sponsor information

Organisation Laxdale Ltd (UK)

Sponsor details

Kings Park House Laurelhill Business Park Polmaise Road Stirling United Kingdom FK7 9JQ +44 (0)1786 476001 admin@Laxdale.co.uk

Sponsor type Industry

Website http://www.laxdale.co.uk

ROR https://ror.org/03gc62f43

Funder(s)

Funder type Industry **Funder Name** Laxdale Ltd (UK)

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 summary Not provided at time of registration