The effects of Sokatin® on mood and cognitive function

| Submission date | Recruitment status | Prospectively registered |
|-------------------|----------------------------------|---|
| 22/10/2010 | No longer recruiting | ☐ Protocol |
| Registration date | Overall study status | Statistical analysis plan |
| 03/12/2010 | Completed | Results |
| Last Edited | Condition category | Individual participant data |
| 03/12/2010 | Mental and Behavioural Disorders | Record updated in last year |

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Dr Esther Boelsma

Contact details

Utrechtseweg 48 Zeist Netherlands 3700

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers

750402.01.028

Study information

Scientific Title

The effects of Sokatin® on mood and cognitive function: a double-blind, placebo-controlled, randomised cross-over study

Study objectives

To investigate if daily oral intake of 500 mg Sokatin® improves mood and cognitive function in healthy subjects.

Ethics approval required

Old ethics approval format

Ethics approval(s)

The Research Subjects and Patients Medical Ethical Review (Medisch-Ethische Toetsing Onderzoek Patienten en Proefpersonen [METOPP]) approved on the 20th October 2010 (ref: M375; NL 33836.028.10)

Study design

Explorative randomised double blind placebo controlled crossover study

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Other

Study type(s)

Treatment

Participant information sheet

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

Health condition(s) or problem(s) studied

Mood/cognitive function

Interventions

Daily intake of one tablet Sokatin® for a period of eight weeks (test) or daily intake of one placebo tablet for a period of eight weeks (control) and vice versa with a wash-out period of 2 weeks in between.

Intervention Type

Drug

Phase

Phase I/II

Drug/device/biological/vaccine name(s)

Sokatin®

Primary outcome measure

- 1. Cognitive performance:
- 1.1. Emotional Stroop Test
- 1.2. Colour Word Vigilance Test
- 1.3. N-back Test
- 1.4. Vigilance and Tracking Test
- 1.5. Switching Attention Test
- 1.6. Long-term Memory Task
- 2. Profile of Mood States Questionnaire

The assessment of mood and cognitive performance using the selected cognitive tests of a computerised validated test system are performed on day 01, day 57, day 71 and day 127.

Secondary outcome measures

No secondary outcome measures

Overall study start date

25/10/2010

Completion date

28/02/2011

Eligibility

Key inclusion criteria

- 1. Healthy volunteers (male and female) aged 30 to 50 years
- 2. Able to perform easy actions on a computer
- 3. Candidates with scores greater than or equal to 45 in the State-Trait Anxiety Inventory (STAI-
- T) during screening
- 4. Having given written informed consent
- 5. Willing to comply with the study procedures

Participant type(s)

Healthy volunteer

Age group

Adult

Sex

Both

Target number of participants

34

Key exclusion criteria

- 1. Participation in any clinical trial up to 90 days before Day 01 of this study
- 2. Participation in any non-invasive clinical trial up to 30 days before Day 01 of this study
- 3. Having a history of medical or surgical events that may significantly affect the study outcome, including psychiatric disorders
- 4. Being colour-blind

- 5. Use of antidepressants
- 6. Being hypersensitive to any ingredient of the study substances
- 7. Use of supplements from screening towards the end of the study
- 8. Being a regular user of recreational drugs
- 9. Excessive alcohol consumption or excessive use of tobacco
- 10. Reported slimming or medically prescribed diet
- 11. Pregnant or lactating or wishing to become pregnant in the period of the study
- 12. Not having a general practitioner

Date of first enrolment

25/10/2010

Date of final enrolment

28/02/2011

Locations

Countries of recruitment

Netherlands

Study participating centre Utrechtseweg 48

Zeist Netherlands 3700

Sponsor information

Organisation

Dr. Willmar Schwabe GmbH & Co. KG (Germany)

Sponsor details

Willmar-Schwabe-Strasse 4 Karlsruhe Germany 76227

Sponsor type

Industry

ROR

https://ror.org/043rrkc78

Funder(s)

Funder type Industry

Funder Name

Dr. Willmar Schwabe GmbH & Co. KG (Germany)

Results and Publications

Publication and dissemination planNot provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

IPD sharing plan summaryNot provided at time of registration