Efficacy and safety of two doses of S 90098 (1 and 2 mg/day), sublingual formulation for 8 weeks in out-patients with major depressive disorder: An 8-week randomised, double-blind, fixed dose, international, multicentre, placebocontrolled study with parallel groups, followed by an extension double-blind treatment period for 16 weeks

Submission date	<b>Recruitment status</b> No longer recruiting	Prospectively registered		
27/03/2008		☐ Protocol		
Registration date 24/04/2008	Overall study status Completed	Statistical analysis plan		
		[X] Results		
Last Edited	Condition category	Individual participant data		
20/04/2020	Mental and Behavioural Disorders			

## Plain English summary of protocol

Not provided at time of registration and not expected to be available in the future

## Contact information

## Type(s)

Scientific

#### Contact name

Prof Frédéric Rouillon

#### Contact details

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## Additional identifiers

#### **EudraCT/CTIS** number

2007-003312-65

**IRAS** number

ClinicalTrials.gov number

#### Secondary identifying numbers

CL2-90098-005

## Study information

#### Scientific Title

Efficacy and safety of two doses of S 90098 (1 and 2 mg/day), sublingual formulation for 8 weeks in out-patients with major depressive disorder: An 8-week randomised, double-blind, fixed dose, international, multicentre, placebo-controlled study with parallel groups, followed by an extension double-blind treatment period for 16 weeks

#### **Study objectives**

To assess the antidepressant efficacy of S 90098 in outpatients suffering from major depressive disorder (MDD).

#### Ethics approval required

Old ethics approval format

## Ethics approval(s)

First ethics committee approval in France on 25/01/2008 (ref: 87-07, CPP Ile de France VI).

## Study design

Randomised, double-blind, parallel group, placebo-controlled, multicentre, phase II study

## 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 the contact details to request a patient information sheet

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

#### Major depressive disorder

#### **Interventions**

Eight-week randomised treatment period with agomelatine orodispersible 1 or 2 mg/day versus placebo followed by an extension double-blind period for 16 weeks.

#### Intervention Type

Drug

#### Phase

Phase II

## Drug/device/biological/vaccine name(s)

S 90098

#### Primary outcome measure

Hamilton Rating Scale for Depression (HAM-D), assessed from baseline to week 24

## Secondary outcome measures

- 1. Safety
- 2. Sleep (Leeds Sleep Evaluation Questionnaire [LSEQ])
- 3. Long term efficacy
- 4. Pharmacokinetic

Outcome measures will be assessed from baseline to week 24.

#### Overall study start date

25/02/2008

## Completion date

30/04/2009

# Eligibility

#### Key inclusion criteria

- 1. Between 18 (or minimum legal age) and 70 years of age
- 2. Out-patients of both genders
- 3. Fulfilling Diagnostic and Statistical Manual of Mental Disorders Fourth Edition (DSM-IV) criteria for major depressive disorder (MDD)

## Participant type(s)

**Patient** 

## Age group

Adult

#### Lower age limit

18 Years

#### Upper age limit

70 Years
Sex Both
Target number of participants 240
Total final enrolment 264
Key exclusion criteria  1. Women of childbearing potential without effective contraception  2. Patients meeting DSM-IV-TR current diagnosis of psychiatric disorder other than MDD  3. Any clinically relevant abnormality detected during the physical examination, electrocardiogram (ECG) or laboratory tests likely to interfere with the study conduct or evaluations
Date of first enrolment 25/02/2008
Date of final enrolment 30/04/2009
Locations
Countries of recruitment Czech Republic
Estonia
Finland
France
Lithuania
Study participating centre

Study participating centre
Centre Hospitalier de Sainte Anne CMME
PARIS cedex 14
France
75674

# Sponsor information

## Organisation

Institut de Recherches Internationales Servier (France)

#### Sponsor details

50 rue Carnot Suresnes France 92284

#### Sponsor type

Industry

#### Website

http://www.servier.com/

#### **ROR**

https://ror.org/034e7c066

# Funder(s)

#### Funder type

Industry

#### Funder Name

Institut de Recherches Internationales Servier (France)

## **Results and Publications**

## Publication and dissemination plan

Publication plan:

Summary results are published in https://clinicaltrials.servier.com.

## Intention to publish date

## Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be available upon request from https://clinicaltrials.servier.com if a Marketing Authorisation has been granted after 1st January 2014.

## IPD sharing plan summary

Available on request

#### **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Basic results				No	No
Basic results			20/04/2020	No	No