# PERSPECTIVES: Protelos® for postmenauposal osteoporotic patients with previous therapies

| Submission date   | Recruitment status  No longer recruiting | <ul><li>Prospectively registered</li><li>Protocol</li></ul> |  |  |
|-------------------|--|---|--|--|
| 20/02/2009        |  |   |  |  |
| Registration date | Overall study status                     | Statistical analysis plan                                   |  |  |
| 15/05/2009        | Completed                                | [X] Results   |  |  |
| Last Edited       | Condition category                       | Individual participant data                                 |  |  |
| 23/01/2019        | Musculoskeletal Diseases                 |   |  |  |

## 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

Dr Martin Kühn

#### Contact details

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# Additional identifiers

#### Protocol serial number

IC4-12911-101-DEU

# Study information

#### Scientific Title

PERSPECTIVES: Protelos® for postmenauposal osteoporotic patients with previous therapies - an observational prospective multi-centre study

#### **Acronym**

**PERSPECTIVES** 

#### **Study objectives**

Effects of Protelos® therapy on osteoporosis and osteoporosis symptoms under practice daily routine in a non-interventional trial.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Ethics Committee of Charité - University Medicine Berlin (Charité - Universitätsmedizin Berlin), Campus Benjamin Franklin, approved on 08/01/2009 (ref: EA4/101/08)

#### Study design

Observational prospective longitudinal multi-centre study

#### Primary study design

Observational

#### Study type(s)

Treatment

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

Postmenopausal osteoporosis

#### **Interventions**

- 1. To get information on the evolution of osteoporosis therapy in the treatment postmenopausal osteoporosis via standardised documentation
- 2. To get information on comobidity and comedication in postmenopausal osteoporosis via standardised documentation.
- 3. Evaluation of osteoporose-associated pain via Visual Analogue Scale (VAS)
- 4. Recording of new osteoporose-associated fractures via standardised documentation
- 5. Evaluation of the efficacy under routine conditions and tolerability of Protelos via Ordinal Scale
- 6. Recording of the osteoporosis-comedications via standardised documentation
- 7. Analysis of the adverse drug reactions of Protelos via standardised adverse drug reactions documentation
- 8. A patients questionnaire (at initiation visit and at final visit) will provide information on:
- 8.1. Evaluation of the quality of life (patients questionnaire including 9 questions = ordinal scale)
- 8.2. Evaluation of the compliance, persistence and adherence with Protelos®
- 8.3. Evaluation of non-medical therapies of postmenopausal osteoporosis (standardised patients questionnaire)

Total duration of follow-up: 3 months

#### Intervention Type

Drug

#### Phase

Phase IV

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

Strontium ranelate (Protelos®)

#### Primary outcome(s)

- 1. Efficacy under routine conditions: ordinal scale (very good, good, moderate, bad)
- 2. Tolerability: ordinal scale (very good, good, moderate, bad)
- 3. Pain (VAS; 0 = no pain 10 = unbearable pain)
- 4. Fractures (standardized documentation)
- 5. Quality of life (patients questionnaire including 9 questions = ordinal scale)
- 6. Compliance, persistence, adherence (standardised patients questionnaire)

All data will be assessed at initiation visit and 3 months later at final visit.

#### Key secondary outcome(s))

No secondary outcome measures

#### Completion date

30/09/2009

# Eligibility

#### Key inclusion criteria

Postmenopausal osteoporotic women, treated with strontium ranelat (Protelos®).

## Participant type(s)

**Patient** 

## Healthy volunteers allowed

No

#### Age group

Adult

#### Sex

**Female** 

#### Key exclusion criteria

Does not meet inclusion criteria

#### Date of first enrolment

02/02/2009

#### Date of final enrolment

30/09/2009

# Locations

#### Countries of recruitment

Germany

Study participating centre Servier Deutschland GmbH Munich Germany 80687

# Sponsor information

## Organisation

Servier Deutschland GmbH (Germany)

#### **ROR**

https://ror.org/05wk4ae67

# Funder(s)

# Funder type

Industry

#### **Funder Name**

Servier Deutschland GmbH (Germany)

# **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                       | 01/01/2012              | Yes            | No              |
| Participant information sheet | Participant information sheet | 11/11/2025 11/11/2025   | 5 No           | Yes             |